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UTILITY PATENT A	PPLICATION TRANSMITT	AL UNDER 37 CFR §1.53(b)
Attorney Docket Number 07891/003005		
Applicant	ROBERT G. KORNELUK STEPHEN BAIRD, AND F	, ALEXANDER E. MACKENZIE, PETER LISTON
Title	MAMMALIAN IAP GENE DETECTION METHODS	FAMILY, PRIMERS, PROBES, and
PRIORITY INFORMATION:		
This application is a continuation 08/576,956, filed December 22, 1 application 08/511,485, filed Augustian	995: which is a Continuation	n-in-Part of United States patent
APPLICATION ELEMENTS:		
Cover sheet		1 page
Specification		88 pages
Claims		6 pages
Abstract		1 page
Drawing		50 sheets
Combined Declaration and POA,  Unsigned; Newly signed for this application A copy from prior application of the disclosure of the prior application of the disclosure of this new application of the disclosure of the third new application of the disclosure of the prior application of the disclosure of the third new application of the disclosure of the third new application of the disclosure of the third new application of the disclosure of the prior application of the disclosure of the third new application of the disclosure of the prior application of the disclosure of	on; 8/576,956 and the entire is considered as being part ication and is hereby	2 pages
Sequence Statement		2 pages
Sequence Listing on Paper		42 pages
Sequence Listing on Diskette		1 diskette
Small Entity Statement, which is:  ☐ Unsigned; ☐ Newly signed for this application; ☐ A copy from prior application 08/576,956 and such small entity status is still proper and desired.		2 pages

Preliminary Amendment	16 pages
IDS	2 pages
Form PTO 1449	5 pages
Cited References	0 references
Recordation Form Cover Sheet and Assignment	0 page
Assignee's Statement	0 page
English Translation	0 page
Certified Copy of Priority Document	0 page
Return Receipt Postcard	1
FILING FEES:	
Basic Filing Fee: \$345	\$345.00
Excess Claims Fee: 47 - 20 = 27 x \$9	\$243.00
Excess Independent Claims Fee: 16 - 3 = 13 x \$39	\$507.00
Multiple Dependent Claims Fee: \$260/\$130	
Total Fees:	\$1095.00
<ul> <li>Enclosed is a check for \$1095.00 to cover the total fees</li> <li>Charge [**AMOUNT**] to Deposit Account No. 03-2095</li> <li>The filing fee is not being paid at this time.</li> <li>Please apply any other charges, or any credits, to Deposit</li> </ul>	to cover the total fees.
CORRESPONDENCE ADDRESS:	
Kristina Bieker-Brady, Ph.D. Reg. No. 39,109 Clark & Elbing LLP 176 Federal Street Boston, MA 02110	Telephone: 617-428-0200 Facsimile: 617-428-7045
Signature James Dk Cany Reg. No. 43.	9/1/00 Date

07891.003005 Utility Application Transmittal Form.wpd



Applicant or Patentee: Robert G. Korneluk et al.

Serial or Patent No.: Filed or Issued: 08/576,956 December 22, 1995

For:

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MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS

## VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 CFR 1.9(f) and 1.27(d)) - NONPROFIT ORGANIZATION

I hereby declare that I am an official empowered to act on behalf of the nonprofit organization identified below:

Name of Organization: University of Ottawa
Address of Organization: 550 Cumberland, Ottawa, Ontario, Canada K1N 6N5
Type of Organization:

[X] UNIVERSITY OR OTHER INSTITUTION OF HIGHER EDUCATION

[ ] TAX EXEMPT UNDER INTERNAL REVENUE SERVICE CODE (26 USC 501(a) and 501(c)(3))

[ ] NONPROFIT SCIENTIFIC OR EDUCATIONAL UNDER STATUTE OF STATE OF THE UNITED STATES OF AMERICA (NAME OF STATE: )

(CITATION OF STATUTE: )

[ ] WOULD QUALIFY AS TAX EXEMPT UNDER INTERNAL REVENUE SERVICE CODE (26 USC 501(a) and 501(c)(3)) IF

LOCATED IN THE UNITED STATES OF AMERICA

[ ] WOULD QUALIFY AS NONPROFIT SCIENTIFIC OR EDUCATIONAL UNDER STATUTE OF STATE OF THE UNITED STATES OF AMERICA (NAME OF STATE: )

(CITATION OF STATUTE: )

I hereby declare that the nonprofit organization identified above qualifies as a nonprofit organization as defined in 37 CFR 19(e) for purposes of paying reduced fees under section 41(a) and (b) of Title 35, United States Code with regard to the invention entitled MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS by inventor(s) Robert G. Korneluk, Atexander R. MacKenzie, and Stephen Baird described in

[] the specification filed herewith.
[X] application serial no. 08/567,959, filed December 22, 1995.

[] patent no. , issued .

hereby declare that rights under contract or law have been conveyed to and remain with the nonprofit organization with regard to the above identified invention.

If the rights held by the nonprofit organization are not exclusive, each individual, concern or organization having rights to the invention are held by any person, other than the inventor, who could not qualify as a small business concern under 37 CFR 1.9(c) or by any concern which would not qualify as a small business concern under 37 CFR 1.9(e).

\*NOTE: Separate verified statements are required from each named person, concern or organization having rights to the invention averring to their status as small entities. (37 CFR 1.27)

Full Name: Apoptogen, Inc.

Address: 100 International Blvd., Etobicoke, Ontario, Canada M9W 6J6

[] INDIVIDUAL [X] SMALL BUSINESS CONCERN [] NONPROFIT ORGANIZATION

I acknowledge the duty to file, in this application or patent, notification of any change in status resulting in loss of entitlement to small entity status prior to paying, or at the time of paying, the earliest of the issue fee or any maintenance fee due after the date on which status as a small entity is no longer appropriate. (37 CFR 1.28(b))

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under section 1001 of Title 18 of the United States Code, and that such willful false statements may jeopardize the validity of the application, any patent issuing thereon, or any patent to which this verified statement is directed.

Name:	Jean Farrall			<del></del>
ritle:	Director, Research Services			
Address:	University of Ottawa, 115 Seraphin Marion, Ottawa, Canada			
Signature:	Jean Farrall	Date:	15 March	1991
				7776

FM Hlbeson

Applicant or Patentee: Robert G. Korneluk et al.

Filed or Issued:

Serial or Patent No.: 08/576,956

December 22, 1995

For:

Signature:

MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTIONS METHODS

#### VERIFIED STATEMENT (DECLARATION) CLAIMING SMALL ENTITY STATUS (37 CFR 1.9(f) and 1.27(c)) - SMALL BUSINESS CONCERN

hereb	y dec	lare	that	I	an
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	[X]		of the small business concern identified below; al of the small business concern empowered to act on behalf of the concern identified
Name of Sm	all Business	Concern:	Apoptogen, Inc.
Address of	Small Busin	ess Concern:	100 International Blvd., Etobicoke, Ontario, Canada M9W 6J6
121.12, and Office, in purposes of of the condiscal year	d reproduced that the number of this state cern of the person, and (2) cerns.	in 37 CFR 1 mber of empl ment, (1) the persons empl oncerns are	entified small business concern qualifies as a small business concern as defined in 13 CFR .9(d), for purposes of paying reduced fees to the United States Patent and Trademark oyees of the concern, including those of its affiliates, does not exceed 500 persons. For e number of employees of the business concern is the average over the previous fiscal year oyed on a full-time, part-time or temporary basis during each of the pay periods of the affiliates of each other when either, directly or indirectly, one concern controls or has a third party or parties controls or has the power to control both.
identified inventor(s)	above with I	regard to the	contract or law have been conveyed to and remain with the small business concern e invention, entitled MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS by exander R. MacKenzie, and Stephen Baird described in
	[X] ap		cion filed herewith. Prial no. 08/576,956, filed December 22, 1995. Ssued .
If the righ having righ inventor, w concern whi 149(e).	its to the ir who would not ch would not OTE: Separat	nvention is to qualify as qualify as to everified s	entified small business concern are not exclusive, each individual, concern or organization listed below and no rights to the invention are held by any person, other than the an independent inventor under 37 CFR 1.9(c) if that person made the invention, or by any a small business concern under 37 CFR 1.9(d), or a nonprofit organization under 37 CFR statements are required from each named person, concern or organization having rights to atus as small entities. (37 CFR 1.27)
Full Name:	University	of Ottawa	
Meddress:	550 Cumberl	and, Ottawa,	Ontario, Canada K1N 6N5
anagii 	[] IN	DIVIDUAL	[ ] SMALL BUSINESS CONCERN [X] NONPROFIT ORGANIZATION
entitlement	to small en	itity status	this application or patent, notification of any change in status resulting in loss of prior to paying, or at the time of paying, the earliest of the issue fee or any on which status as a small entity is no longer appropriate. (37 CFR 1.28(b))
pelief are and the lik Code, and t	believed to e so made ar hat such wil	be true; and e punishable lful false s	is made herein of my own knowledge are true and that all statements made on information and if further that these statements were made with the knowledge that willful false statements by fine or imprisonment, or both, under section 1001 of Title 18 of the United States statements may jeopardize the validity of the application, any patent issuing thereon, or statement is directed.
lame:	Frank Glees	on	
itle:	President a	nd CEO	·
ddreec-	180 Etchico	ka Ontario	Canada MOU 616 100 International Stud

Date: \_\_

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## IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: Robert G. Korneluk et al.

Art Unit:

Not Yet Assigned

Serial No.:

Not Yet Assigned

Examiner:

Not Yet Assigned

Filed:

September 1, 2000

Title:

MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND

**DETECTION METHODS** 

Director for Patents Washington, D.C. 20231

### PRELIMINARY AMENDMENT

Prior to examination of the above-referenced application, kindly consider the following amendments and remarks.

Please amend the application as follows:

### In the specification:

At page 1, line 5, before "The invention relates to apoptosis.", add the following:

## -- Cross Reference To Related Applications

This application is a continuation of U.S.S.N. 08/576,956, filed December 22, 1995, which is a continuation-in-part of U.S.S.N. 08/511,485, filed August 4, 1995, now issued as U.S. Patent No. 5,919,912.--.

At page 6, line 27, replace "IAP disease resistance gene" with --IAP gene--.

At page 18, line 15, replace "Fig. 10 is a Northern blot" with -- Figs. 10A-C are a series of Northern blots --.

At page 18, line 17, replace "Fig. 11 is a Northern blot" with -- Figs. 11A-C are a series of Northern blots --.

At page 18, line 19, replace "Fig. 12 is a Northern blot" with -- Figs. 12A-C are a series of Northern blots --.

At page 19, line 1, after "Tables 1 and 2"insert --(SEQ ID NOS: 45-92)--.

At page 24, line 23, after "MEQKLISEEDL," insert -- (SEQ ID NO: 43) --.

At page 26, line 23, replace "Embo," with -- EMBO --.

At page 27, line 3, replace "Neurobiol," with -- Neurobiol. --.

At page 27, line 27, replace "Virol," with -- Virol. --.

At page 34, line 18, replace "Cell," with -- Cell --.

At page 34, line 18, replace "Nature," with -- Nature --.

At page 36, line 8, replace "Med," with -- Med. --.

Kindly remove the sequence listing found at pages 51-88 and renumber the pages

of the claims and abstract consecutively thereafter. The enclosed amended sequence listing should be inserted at the end of the application.

#### In the Claims:

Cancel claims 2, 15-29, and 33-47.

Amend claims 1, 3-7, 13, 14, and 30-32 as follows.

- 1. (Amended) A substantially [Substantially] pure nucleic acid encoding [an IAP] a mammalian inhibitor of apoptosis protein (IAP) polypeptide, wherein said inhibitor of apoptosis protein is a protein that modulates apoptosis and comprises a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of apoptosis repeat (BIR) domain.
- 3. (Amended) The nucleic acid of claim [2] 1, wherein said polypeptide has at least two [BIR] baculovirus inhibitor of apoptosis repeat (BIR) domains.
- 4. (Amended) The nucleic acid of claim 3, wherein said polypeptide has at least three [BIR] <u>baculovirus inhibitor of apoptosis repeat (BIR)</u> domains.
- 5. (Amended) The nucleic acid of claim 1, wherein said [DNA] <u>nucleic acid</u> contains the [xiap] <u>X-linked inhibitor of apoptosis protein (xiap)</u> gene.

- 6. (Amended) The nucleic acid of claim 1, wherein said [DNA] <u>nucleic acid</u> contains the [hiap2] <u>human inhibitor of apoptosis protein 2 (hiap2)</u> gene.
- 7. (Amended) The nucleic acid of claim 1, wherein said [DNA] <u>nucleic acid</u> contains the [hiap1] <u>human inhibitor of apoptosis protein 1 (hiap1)</u> gene.

  DNA.
- 13. (Amended) A [Substantially] substantially pure [DNA] nucleic acid having the sequence of Fig. 5 (SEQ ID NO: 39), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 5 (SEQ ID NO: 40).
- 14. (Amended) A [Substantially] substantially pure [DNA] nucleic acid having the sequence of Fig. 6 (SEQ ID NO: 41), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 6 (SEQ ID NO: 42).
- 30. (Amended) A method of producing [an IAP] <u>a mammalian inhibitor of apoptosis protein (IAP)</u> polypeptide comprising:

providing a cell transformed with [DNA] <u>nucleic acid</u> encoding [an] <u>a mammalian</u>

IAP polypeptide positioned for expression in said cell, <u>said polypeptide comprising a ring</u>

<u>zinc finger (RZF) domain;</u>

culturing said transformed cell under conditions for expressing said [DNA] <u>nucleic</u> <u>acid</u>; and

[isolating] producing said IAP polypeptide.

- 31. (Amended) The method of claim 30, wherein said <u>mammalian inhibitor of</u> apoptosis (IAP) [IAP] polypeptide is murine <u>human inhibitor of apoptosis protein 1 (m-HIAP1)</u> [HIAP1].
- 32. (Amended) The method of claim 30, wherein said <u>mammalian inhibitor of</u> apoptosis (IAP) [IAP] polypeptide is murine <u>human inhibitor of apoptosis protein 2 (m-HIAP2)</u> [HIAP2].

Add the following new claims 48-78.

--48. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the nucleic acid sequence of Fig. 5 (SEQ ID NO: 39), wherein said nucleic acid hybridizes to said probe under low stringency conditions, said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus

inhibitor of apoptosis repeat (BIR) domain.

- 49. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the nucleic acid sequence of Fig. 6 (SEQ ID NO: 41), wherein said nucleic acid hybridizes to said probe under low stringency conditions, said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of apoptosis repeat (BIR) domain.
- 50. A substantially pure nucleic acid encoding a baculovirus inhibitor of apoptosis repeat (BIR) domain, said nucleic acid comprising a sequence selected from the group consisting of SEQ ID NO: 45, SEQ ID NO: 46, SEQ ID NO: 47, SEQ ID NO: 49, SEQ ID NO: 50, SEQ ID NO: 51, SEQ ID NO: 53, SEQ ID NO: 54, SEQ ID NO: 55, SEQ ID NO: 57, SEQ ID NO: 58, SEQ ID NO: 59, SEQ ID NO: 61, SEQ ID NO: 62, SEQ ID NO: 63, SEQ ID NO: 65, SEQ ID NO: 66, and SEQ ID NO: 67.
- 51. A substantially pure nucleic acid encoding a ring zinc finger (RZF) domain, said nucleic acid comprising a sequence selected from the group consisting of SEQ ID NO: 48, SEQ ID NO: 52, SEQ ID NO: 56, SEQ ID NO: 60, SEQ ID NO: 64, and SEQ ID

- 52. The nucleic acid of claim 1, wherein said nucleic acid encodes an X-linked inhibitor of apoptosis protein (XIAP).
- 53. The nucleic acid of claim 52, wherein said X-linked inhibitor of apoptosis protein (XIAP) is from a mouse.
- 54. The nucleic acid of claim 52, wherein said X-linked inhibitor of apoptosis protein (XIAP) is from a human.
- 55. The nucleic acid of claim 1, wherein said nucleic acid encodes a human inhibitor of apoptosis protein 1 (HIAP1).
- 56. The nucleic acid of claim 55, wherein said human inhibitor of apoptosis protein 1 (HIAP1) is from a mouse.
- 57. The nucleic acid of claim 55, wherein said human inhibitor of apoptosis protein 1 (HIAP1) is from a human.

- 58. The nucleic acid of claim 1, wherein said nucleic acid encodes a human inhibitor of apoptosis protein 2 (HIAP2).
- 59. The nucleic acid of claim 58, wherein said human inhibitor of apoptosis protein 2 (HIAP2) is from a mouse.
- 60. The nucleic acid of claim 58, wherein said human inhibitor of apoptosis protein 2 (HIAP2) is from a human.
- 61. The nucleic acid of claim 5, wherein said X-linked inhibitor of apoptosis protein (xiap) gene is from a mouse.
- 62. The nucleic acid of claim 5, wherein said X-linked inhibitor of apoptosis protein (xiap) gene is from a human.
- 63. The nucleic acid of claim 6, wherein said human inhibitor of apoptosis protein 2 (hiap2) gene is from a mouse.
- 64. The nucleic acid of claim 6, wherein said human inhibitor of apoptosis protein 2 (hiap2) gene is from a human.

- 65. The nucleic acid of claim 7, wherein said human inhibitor of apoptosis protein 1 (hiap1) gene is from a mouse.
- 66. The nucleic acid of claim 7, wherein said human inhibitor of apoptosis protein 1 (hiap1) gene is from a human.
- 67. A substantially pure nucleic acid having the sequence of Fig. 1 (SEQ ID NO: 3), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 1 (SEQ ID NO: 4).
- 68. A substantially pure nucleic acid having the sequence of Fig. 2 (SEQ ID NO: 5), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 2 (SEQ ID NO: 6).
- 69. A substantially pure nucleic acid having the sequence of Fig. 3 (SEQ ID NO: 7), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 3 (SEQ ID NO: 8).
- 70. A substantially pure nucleic acid having the sequence of Fig. 4 (SEQ ID NO: 9), or degenerate variants thereof, and encoding the amino acid sequence of Fig. 4 (SEQ

- 71. The method of claim 30, wherein said mammalian inhibitor of apoptosis protein (IAP) polypeptide is human inhibitor of apoptosis protein 1 (HIAP1).
- 72. The method of claim 30, wherein said mammalian inhibitor of apoptosis protein (IAP) polypeptide is human inhibitor of apoptosis protein 2 (HIAP2).
- 73. The method of claim 30, wherein said mammalian inhibitor of apoptosis protein (IAP) polypeptide is murine X-linked inhibitor of apoptosis protein (m-XIAP).
- 74. The method of claim 30, wherein said mammalian inhibitor of apoptosis protein (IAP) polypeptide is human X-linked inhibitor of apoptosis protein (XIAP).
- 75. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the DNA sequence of Fig. 1 (SEQ ID NO: 3), wherein said nucleic acid hybridizes to said probe under low stringency conditions, said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of

apoptosis repeat (BIR) domain.

- 76. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the DNA sequence of Fig. 2 (SEQ ID NO: 5), wherein said nucleic acid hybridizes to said probe under low stringency conditions, said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of apoptosis repeat (BIR) domain.
- 77. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the DNA sequence of Fig. 3 (SEQ ID NO: 7), wherein said nucleic acid hybridizes to said probe under low stringency conditions, said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of apoptosis repeat (BIR) domain.
- 78. A substantially pure nucleic acid that hybridizes to a probe of at least 40 nucleotides in length, said probe derived from the DNA sequence of Fig. 4 (SEQ ID NO: 9), wherein said nucleic acid hybridizes to said probe under low stringency conditions,

said conditions comprising washing with 2X SSC at 40°C, and wherein said nucleic acid encodes a mammalian inhibitor of apoptosis protein (IAP) polypeptide, said polypeptide comprising a ring zinc finger (RZF) domain and at least one baculovirus inhibitor of apoptosis repeat (BIR) domain.--

#### **REMARKS**

In general, Applicants' presently claimed invention features substantially pure nucleic acids encoding mammalian IAP polypeptides and methods of using such nucleic acids to produce such mammalian IAP polypeptides.

#### Support for the Amendments

The specification and drawings have been amended to comply with the requirements of 37 C.F.R. § 1.821 through 1.825. The specification has also been amended to properly refer to each individual panel of a drawing.

The specification and the claims have been amended to correct regrettable typographical errors.

Applicants have added new claims 48 and 49 to cover substantially pure DNA encoding mammalian inhibitor of apoptosis protein (IAP) polypeptides that hybridize under low stringency conditions to SEQ ID NO: 39 and SEQ ID NO: 41, respectively. Support for these new claims may be found in the specification at page 48, lines 15-20.

Applicants have added new claim 50 to cover a substantially pure DNA encoding a baculovirus inhibitor of apoptosis repeat domain that comprises the sequence of SEQ ID

NO: 45, SEQ ID NO: 46, SEQ ID NO: 47, SEQ ID NO: 49, SEQ ID NO: 50, SEQ ID NO: 51, SEQ ID NO: 53, SEQ ID NO: 54, SEQ ID NO: 55, SEQ ID NO: 57, SEQ ID NO: 58, SEQ ID NO: 59, SEQ ID NO: 61, SEQ ID NO: 62, SEQ ID NO: 63, SEQ ID NO: 65, SEQ ID NO: 66, or SEQ ID NO: 67. Support for this new claim can be found in the specification at page 19 in Table I (page 19, lines 12-20). The DNA of this claim finds use as, for example, a hybridization probe for screening libraries.

Applicants have added new claim 51 to cover a substantially pure DNA encoding a ring zinc finger domain that comprises the sequence of SEQ ID NO: 48, SEQ ID NO: 52, SEQ ID NO: 56, SEQ ID NO: 60, SEQ ID NO: 64, or SEQ ID NO: 68. Support for this new claim can be found in the specification at page 19 in Table I (page 19, lines 12-20). The DNA of this claim finds use as, for example, a hybridization probe for screening libraries.

Applicants have added new claim 52 to claim nucleic acid encoding an X-linked inhibitor of apoptosis protein (XIAP). New dependent claims 53 and 54 have been added to specifically claim nucleic acids encoding XIAP from a mouse and from a human, respectively. Support for these new claims may be found in the specification at page 21, lines 2-21, page 22, lines 8-32, and in Figs. 1 and 4.

Applicants have added new claim 55 to claim nucleic acid encoding a human inhibitor of apoptosis protein 1 (HIAP1). New dependent claims 56 and 57 have been added to specifically claim nucleic acids encoding HIAP1 from a mouse and from a human, respectively. Support for these new claims may be found in the specification at

page 21, line 22 through page 22, line 7, and in Figs. 2 and 5.

Applicants have added new claim 58 to claim nucleic acid encoding a human inhibitor of apoptosis protein 2 (HIAP2). New dependent claims 59 and 60 have been added to specifically claim nucleic acids encoding HIAP2 from a mouse and from a human, respectively. Support for these new claims may be found in the specification at page 21, line 22 through page 22, line 7, and in Figs. 3 and 6.

Applicants have added new dependent claims 61 and 62 to specifically claim nucleic acids containing the X-linked inhibitor of apoptosis (xiap) gene, where the (xiap) gene is from a mouse or a human, respectively. Support for these new claims may be found in the specification at page 21, lines 2-21, page 22, lines 8-32, and in Figs. 1 and 4.

Applicants have added new dependent claims 63 and 64 to specifically claim nucleic acids containing the human inhibitor of apoptosis 2 (hiap2) gene, where the (hiap2) gene is from a mouse or a human, respectively. Support for these new claims may be found in the specification at page 21, line 22 through page 22, line 7, and in Figs. 3 and 6.

Applicants have added new dependent claims 65 and 66 to specifically claim nucleic acids containing the human inhibitor of apoptosis 1 (hiap1) gene, where the (hiap1) gene is from a mouse or a human, respectively. Support for these new claims may be found in the specification at page 21, line 22 through page 22, line 7, and in Figs. 2 and 5.

Applicants have added new claims 67, 68, 69, and 70 to specifically claim substantially pure nucleic acids having the sequence of and encoding the amino acid sequence of Figs. 1, 2, 3, and 4, respectively. Support for these new claims may be found in the specification, for example, at page 21, line 2 through page 22, line 32, and in Figs. 1-4.

Applicants have added new dependent claims 71, 72, 73, and 74 to specifically claim methods for producing human inhibitor of apoptosis protein 1, human inhibitor of apoptosis protein 2, murine X-linked inhibitor of apoptosis protein, and human X-linked inhibitor of apoptosis protein, respectively. Support for these new claims may be found in the specification, for example, at page 5, lines 7-12; at page 21, line 2 through page 22, line 32; and in Figs. 1-4.

Applicants have added new claims 75-78 to specifically claim substantially pure nucleic acids encoding mammalian inhibitor of apoptosis protein (IAP) polypeptides that hybridize under low stringency conditions to probes derived from the DNA sequences of SEQ ID NO: 3, SEQ ID NO: 5, SEQ ID NO: 7, and SEQ ID NO: 9, respectively. Support for these new claims may be found in the specification at page 48, lines 15-20, and in Figs. 1-4. No new matter is added by any of these amendments.

#### Sequence Listing

As required by 37 CFR 1.825(a), enclosed is an amended sequence listing consisting of 42 sheets to be inserted at the end of the application. The amendments to

the sequence listing provide each sequence in the specification with a unique SEQ ID NO, and contain no new matter. In particular, SEQ ID NOS: 69-92 have been added to include the sequences described in Table 2, found at page 20 of the specification.

As required by 37 CFR 1.825(b), also enclosed is a diskette containing a copy of the sequence listing in computer readable form including all previously submitted data with the amendments incorporated therein. The contents of the computer readable form are the same as the contents of the paper sheets.

If there are any charges or any credits, please apply them to Deposit Account No. 03-2095.

Respectfully submitted,

Date:  $\frac{9/1/00}{}$ 

Kristina Bieker-Brady, Ph/D.

Reg. No. 39,109 Tames De Camp Reg. No. 43580

Clark & Elbing LLP 176 Federal Street Boston, MA 02110

Telephone: 617-428-0200

Facsimile: 617-428-7045

07891.003005 Preliminary amendment xxx.wpd

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Date of Deposit <u>September 1, 2000</u> Label No	umber: <u>EL509049835US</u>			
I hereby certify under 37 CFR 1.10 that this correspondence is be "Express Mail Post Office to Addressee" with sufficient postag PATENT APPLICATION, Director for Patents, Washington, D.C.  Luis A. Cruz Printed name of person mailing correspondence	e on the date indicated above and is addressed to BOX			

## **APPLICATION**

## **FOR**

## UNITED STATES LETTERS PATENT

APPLICANT : ROBERT G. KORNELUK, ALEXANDER E. MACKENZIE,

STEPHEN BAIRD, AND PETER LISTON

TITLE : MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES,

AND DETECTION METHODS

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# MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS

The invention relates to apoptosis.

#### Background of the Invention

There are two general ways by which cells die. easily recognized pathway is necrosis, a process of cell death usually resulting from severe and sudden injury. necrosis, changes in cellular homeostasis occur with loss of membrane integrity. Dysregulation of osmotic pressure results and, as a consequence, the cells swell and finally rupture. The cellular contents are then spilled into the surrounding tissue space and, usually, an inflammation response ensues. A second form of cell death is apoptosis. This cell "suicide" pathway or programmed cell death often occurs so rapidly that in some biological systems the apoptotic process is difficult to ascertain. Indeed, it has been only in the past few years that the involvement of apoptosis in a wide spectrum of biological processes has become recognized. Apoptosis is a fundamental physiological pathway of cell death, highly conserved throughout evolution, and playing a major role in development, viral pathogenesis, cancer, autoimmune diseases and neurodegenerative disorders.

Inappropriate increases in apoptosis may cause or contribute to a variety of diseases, including AIDS, neurodegenerative diseases (e.g. Alzheimer's Disease, Parkinson's Disease, Amyotrophic Lateral Sclerosis (ALS), retinitis pigmentosa and other diseases of the retina, myelodysplastic syndrome (e.g., aplastic anemia), toxin-

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induced liver disease (e.g., alcoholism) and ischemic injury (e.g., myocardial infarction, stroke, and reperfusion injury). In addition, disruption of normally occurring apoptosis has been implicated in the development of some cancers (e.g. follicular lymphoma, p53 carcinomas, and hormone dependent tumors), autoimmune disorders (e.g., lupus erythematosis and multiple sclerosis) and viral infections (e.g., herpes virus, poxvirus, and adenovirus infections).

Mature CD4<sup>+</sup> T-lymphocytes in patients with HIV-1 have been observed to respond to stimulation with mitogens or super-antigens by undergoing increased apoptosis. The great majority of these cells are not infected and similar inappropriate antigen-induced apoptosis could be very important in the destruction of this vital part of the immune system early in HIV infection.

Baculoviruses encode <u>i</u>nhibitors of <u>a</u>poptosis proteins (IAPs). These proteins inhibit the apoptosis which otherwise occurs when insect cells are infected by the virus. Baculovirus IAP proteins work in a manner which is thought to be independent of other viral proteins. The baculovirus IAP genes include sequences encoding a ring zinc finger-like motif which is presumed to be involved in the direct binding of DNA.

### Summary of the Invention

In general, the invention features substantially pure DNA (for example, genomic DNA, cDNA, or synthetic DNA) encoding a mammalian IAP polypeptide as defined below. In related aspects, the invention also features a vector, a cell (e.g., a mammalian, yeast or bacterial cell), and a transgenic animal or embryo thereof which includes such a substantially pure DNA encoding an IAP polypeptide.

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In preferred embodiments, an IAP gene is the xiap (including human xiap and its murine homolog, m-xiap), hiap1 (including human hiap1 and m-hiap1), or the hiap2 gene (including human hiap2 and m-hiap2). In most preferred embodiments the IAP gene is a human IAP gene. In other various preferred embodiments, the cell is a transformed cell. In related aspects, the invention features a transgenic animal containing a transgene which encodes an IAP polypeptide that is expressed in or delivered to tissue normally susceptible to apoptosis.

In yet another aspect, the invention features DNA encoding fragments of IAP polypeptides including the BIR domains and the RZF domains provided herein.

In specific embodiments, the invention features DNA sequences substantially identical to the DNA sequences shown in Figs. 1-6.

In another aspect, the invention also features RNA which is encoded by the DNA described herein. Preferably, the RNA is mRNA. In another embodiment the RNA is antisense RNA.

In another aspect, the invention features a substantially pure polypeptide having a sequence substantially identical to one of the IAP amino acid sequences shown in Figures 1-6.

In a second aspect, the invention features a substantially pure DNA which includes a promoter capable of expressing the IAP gene in a cell susceptible to apoptosis. In preferred embodiments, the IAP gene is xiap (including the human or murine xiap), hiapl (preferably the human or murine hiapl), or hiap2 (preferably the human or murine hiap2). hiap2 may be the full length gene, as shown in Fig. 3, or the truncated variant having the sequence boxed in Fig. 3 deleted.

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In preferred embodiments, the promoter is the promoter native to an IAP gene. Additionally, transcriptional and translational regulatory regions are preferably native to an IAP gene.

In another aspect, the invention provides transgenic cell lines and transgenic animals. The transgenic cells of the invention are preferably cells which are susceptible to apoptosis. In preferred embodiments, the transgenic cell is a fibroblast, neuronal cell, a lymphocyte cell, or an insect cell. Most preferably, the neuron is a motor neuron and the lymphocyte is a CD4<sup>+</sup> T-cell.

In another aspect, the invention features a method of inhibiting apoptosis which involves producing a transgenic cell having a transgene encoding an IAP polypeptide wherein the transgene is integrated into the genome of the cell and is positioned for expression in the cell and wherein the IAP transgene is expressed in the cell at a level sufficient to inhibit apoptosis.

In a related aspect, the invention features a transgenic animal, preferably a mammal, more preferably a rodent, and most preferably a mouse, having either increased copies of IAP genes inserted into the genome or a knockout of an IAP gene in the genome. The transgenic animals may express an increased amount of IAP polypeptide or may express a decreased amount of an IAP polypeptide, respectively. In related embodiments, the invention provides a method of utilizing the IAP nucleic acid to engineer a knockout mutation in an IAP gene and a method of making an animal with increased expression by insertion of IAP gene into the genome.

In another aspect, the invention features a method of detecting an IAP in a cell involving: (a) contacting the IAP gene or a portion thereof greater than 9 nucleic acids,

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preferably greater than 18 nucleic acids in length with a preparation of genomic DNA from the cell under hybridization conditions providing detection of DNA sequences having about 50% or greater nucleotide sequence identity to the amino acid encoding DNA sequences of hiap1, hiap2, or xiap IAP polypeptides.

In another aspect, the invention features a method of producing an IAP polypeptide which involves: (a) providing a cell transformed with DNA encoding an IAP polypeptide positioned for expression in the cell; (b) culturing the cell under conditions for expressing the DNA; and (c) isolating the IAP polypeptide. In preferred embodiments the IAP polypeptide is expressed by DNA which has a constituative or inducible promotor. In our embodiment, the promotor is a heterologous promotor.

In another aspect, the invention features substantially pure mammalian IAP polypeptide. Preferably, the polypeptide includes a greater than 50 amino acid sequence substantially identical to a greater than 50 amino acid sequence shown in any one of Figs. 1-4. Most preferably, the polypeptide is the human or murine XIAP, HIAP1, or HIAP2 polypeptide. Fragments including BIR domains and RZF-domains provided herein are also a part of the invention.

In another aspect, the invention features a recombinant mammalian polypeptide capable of modulating apoptosis wherein the polypeptide includes at least a ring zinc finger domain and a BIR domain as defined herein. In preferred embodiments, the invention features a substantially pure polypeptide and an oligonucleotide encoding said polypeptide, the polypeptide including a ring zinc finger (RZF) having the sequence:

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In various preferred embodiments the protein has at least two or, more preferably at least three BIR domains, the RZF domain has one of the IAP sequences shown in Fig. 6, and the BIR domains are comprised of BIR domains shown in Fig. 5. In other preferred embodiments the BIR domains are at the amino terminal end of the protein relative to the RZF domain, which is at or near the carboxy terminus of the polypeptide.

In another aspect, the invention features an IAP gene isolated according to the method involving: (a) providing a sample of DNA; (b) providing a pair of oligonucleotides having sequence homology to a conserved region of an IAP disease-resistance gene; (c) combining the pair of oligonucleotides with the cell DNA sample under conditions suitable for polymerase chain reaction-mediated DNA amplification; and (d) isolating the amplified IAP gene or fragment thereof.

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In preferred embodiments, the amplification is carried out using a reverse-transcription polymerase chain reaction, for example, the RACE method.

In another aspect, the invention features an IAP gene isolated according to the method involving: (a) providing a preparation of DNA; (b) providing a detectably-labelled DNA sequence having homology to a conserved region of an IAP gene; (c) contacting the preparation of DNA with the detectably-labelled DNA sequence under hybridization conditions providing detection of genes having 50% or greater nucleotide sequence identity; and (d) identifying an IAP gene by its association with the detectable label.

In another aspect, the invention features an IAP gene isolated according to the method involving: (a) providing a cell sample; (b) introducing by transformation into the cell sample a candidate IAP gene; (c) expressing the candidate IAP gene within the cell sample; and (d) determining whether the cell sample exhibits an altered apoptotic response, whereby a response identifies an IAP gene.

In another aspect, the invention features a method of identifying an TAP gene in a cell, involving: (a) providing a preparation of cellular DNA (for example, from the human genome or a cDNA library (such as a cDNA library isolated from a cell type which undergoes apoptosis); (b) providing a detectably-labelled DNA sequence (for example, prepared by the methods of the invention) having homology to a conserved region of an TAP gene; (c) contacting the preparation of cellular DNA with the detectably-labelled DNA sequence under hybridization conditions providing detection of genes having 50% nucleotide or greater sequence identity; and (d) identifying an TAP gene by its association with the detectable label.

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In another aspect, the invention features a method of isolating an IAP gene from a recombinant library, involving: (a) providing a recombinant library; (b) contacting the library with a detectably-labelled gene fragment produced according to the PCR method of the invention under hybridization conditions providing detection of genes having 50% or greater nucleotide sequence identity; and (c) isolating an IAP gene by its association with the detectable label.

In another aspect, the invention features a method of identifying an IAP gene involving: (a) providing a cell tissue sample; (b) introducing by transformation into the cell sample a candidate IAP gene; (c) expressing the candidate IAP gene within the cell sample; and (d) determining whether the cell sample exhibits inhibition of apoptosis, whereby a change in (i.e. modulation of) apoptosis identifies an IAP gene.

Preferably, the cell sample is a cell type which may be assayed for apoptosis (e.g., lymphocytes, T-cells and B-cells, neuronal cells, baculovirus infected insect cells and fibroblast cells); the candidate IAP gene is obtained from a cDNA expression library; and the apoptosis response is the inhibition of apoptosis.

In another aspect, the invention features a method of inhibiting apoptosis in a mammal wherein the method includes: (a) providing DNA encoding at least one IAP polypeptide to a cell which is susceptible to apoptosis; wherein the DNA is integrated into the genome of the cell and is positioned for expression in the cell; and the IAP gene is under the control of regulatory sequences suitable for controlled expression of the gene(s); wherein the IAP transgene is expressed at a level sufficient to inhibit apoptosis relative to a cell lacking the IAP transgene. It

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will be appreciated that IAP polypeptides also may be administered directly to inhibit any undesirable apoptosis.

In a related aspect, the invention features a method of inhibiting apoptosis wherein the method involves: (a) producing a cell having integrated in the genome a transgene containing the IAP gene under the control of a promoter providing constitutive expression of the IAP gene.

In yet another related aspect, the invention features a method of inhibiting apoptosis wherein the method involves: (a) producing a cell having integrated in the genome a transgene containing the IAP gene under the control of a promoter providing controllable expression of the IAP gene; and (b) regulating the environment of the cell so that the IAP transgene is controllably expressed in the cell. preferred embodiments, the IAP gene is expressed using a tissue-specific or cell type-specific promoter, or by a promoter that is activated by the introduction of an external signal or agent, such as a chemical signal or agent. In preferred embodiments the cell is a lymphocyte or B-cell, a neuronal cell, or a fibroblast. In other embodiments the cell is a cell in an HIV infected human, or a mammal with a neurodegenerative disease, ischemia, toxin induced liver disease, or a myelodysplastic syndrome.

In a related aspect, the invention provides a method of inhibiting apoptosis in a mammal by providing an apoptosis-inhibiting amount of IAP polypeptide.

In another aspect, the invention features a purified antibody which binds specifically to an IAP family protein. Such an antibody may be used in any standard immunodetection method for the identification of an IAP polypeptide. Preferably, the antibody binds specifically to xiap, hiap1 or hiap2. In various embodiments the antibody may react with other IAP polypeptides or may be specific for one or a

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few IAP polypeptides. The antibody may be a monoclonal polyclonal antibody. Preferably, the antibody reacts specifically with only one of the IAP polypeptides, for example, reacts with murine and human xiap, but not with hiap1 or hiap2 from mammalian species.

In another aspect, the invention features a method of identifying a compound which modulates apoptosis. The method includes (a) providing a cell expressing an IAP polypeptide; and (b) contracting the cell with a candidate compound, and monitoring the expression of an IAP gene. An alteration in the level of expression of the IAP gene indicates the presence of a compound which modulates apoptosis. The compound may be an inhibitor or an enhancer of apoptosis. In various preferred embodiments, the cell is a fibroblast, a neuronal cell, a lymphocyte (T-cell or B-cell), or an insect cell; the polypeptide expression being monitored is XIAP, HIAP1, or HIAP2 (e.g., human or murine).

In a related aspect, the invention features methods of detecting compounds which modulate apoptosis using the interaction trap technology and IAP polypeptides or fragments thereof as a component of the bait. In preferred embodiments, the compound being tested as a modulator of apoptosis is also a polypeptide.

In another aspect, the invention features a method for diagnosing a cell proliferation disease, or an increased liklihood of such a disease, using an IAP nucleic acid probe or antibody. Preferably, the disease is a cancer. Most preferably, the disease is selected from the group consisting of promyelocytic leukemia, a Hela-type carcinoma, chronic myelogenous leukemia (preferably using xiap or hiap2 related probes), lymphoblastic leukemia (preferably using a xiap related probe), Burkitt's lymphoma (preferably using an hiap1 related probe), colorectal adenocarcinoma, lung

carcinoma, and melanoma (preferably using a xiap probe). Preferably, a diagnosis is indicated by a 2-fold increase in expression or activity, more preferably, at least a 10-fold increase in expression or activity.

By "IAP gene" is meant a gene encoding a polypeptide having at least one BIR domain and a ring zinc finger domain which is capable of modulating (inhibiting or enhancing) apoptosis in a cell or tissue when provided by other intracellular or extracellular delivery methods. In preferred embodiments the IAP gene is a gene having about 50% or greater nucleotide sequence identity to at least one of the IAP amino acid encoding sequences of Figs. 1-4 or portions thereof. Preferably, the region of sequence over which identity is measured is a region encoding at least one BIR domain and a ring zinc finger domain. Mammalian IAP genes include nucleotide sequences isolated from any mammalian source. Preferably, the mammal is a human.

By an "IAP gene" is also meant any member of the family of apoptosis inhibitory genes characterized by their ability to modulate apoptosis and having at least 20%, preferably 30%, and most preferably 50% amino acid sequence identity to at least one of the conserved regions of one of the IAP members described herein (i.e., either the BIR or ring zinc finger domains from the human or murine xiap, hiap1 and hiap2). Representative members of the IAP gene family include, without limitation, the human and murine xiap, hiap1, and hiap2 genes. By "IAP protein" is meant a polypeptide encoded by an IAP gene.

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By "modulating apoptosis" or "altering apoptosis" is meant increasing or decreasing the number of cells which undergo apoptosis in a given cell population. Preferably, the cell population is selected from a group including T-cells, neuronal cells, fibroblasts, or any other cell line known to undergo apoptosis in a laboratory setting (e.g., the baculovirus infected insect cells). It will be appreciated that the degree of modulation provided by an IAP or modulating compound in a given assay will vary, but that one skilled in the art can determine the statistically significant change in the level of apoptosis which identifies an IAP or a compound which modulates an IAP.

By "inhibiting apoptosis" is meant any decrease in the number of cells which undergo apoptosis relative to an untreated control. Preferably, the decrease is at least 25%, more preferably the decrease is 50%, and most preferably the decrease is at least one-fold.

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By "polypeptide" is meant any chain of amino acids, regardless of length or post-translational modification (e.g., glycosylation or phosphorylation).

By "substantially identical" is meant a polypeptide or nucleic acid exhibiting at least 50%, preferably 85%, more preferably 90%, and most preferably 95% homology to a reference amino acid or nucleic acid sequence. For polypeptides, the length of comparison sequences will generally be at least 16 amino acids, preferably at least 20 amino acids, more preferably at least 25 amino acids, and most preferably 35 amino acids. For nucleic acids, the length of comparison sequences will generally be at least 50 nucleotides, preferably at least 60 nucleotides, more preferably at least 75 nucleotides, and most preferably 110 nucleotides.

Sequence identity is typically measured using sequence analysis software (e.g., Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin Biotechnology Center, 1710 University Avenue, Madison, WI 53705). Such software matches similar sequences by assigning degrees of homology to various substitutions, deletions, and other modifications. Conservative substitutions typically include substitutions within the following groups: glycine, alanine, valine, isoleucine, leucine; aspartic acid, glutamic acid, asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine.

By a "substantially pure polypeptide" is meant an IAP polypeptide which has been separated from components which naturally accompany it. Typically, the polypeptide is substantially pure when it is at least 60%, by weight, free from the proteins and naturally-occurring organic molecules with which it is naturally associated. Preferably, the

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preparation is at least 75%, more preferably at least 90%, and most preferably at least 99%, by weight, IAP polypeptide. A substantially pure IAP polypeptide may be obtained, for example, by extraction from a natural source (e.g., a fibroblast, neuronal cell, or lymphocyte cell); by expression of a recombinant nucleic acid encoding an IAP polypeptide; or by chemically synthesizing the protein. Purity can be measured by any appropriate method, e.g., those described in column chromatography, polyacrylamide gel electrophoresis, or by HPLC analysis.

A protein is substantially free of naturally associated components when it is separated from those contaminants which accompany it in its natural state. Thus, a protein which is chemically synthesized or produced in a cellular system different from the cell from which it naturally originates will be substantially free from its naturally associated components. Accordingly, substantially pure polypeptides include those derived from eukaryotic organisms but synthesized in *E. coli* or other prokaryotes.

By "substantially pure DNA" is meant DNA that is free of the genes which, in the naturally-occurring genome of the organism from which the DNA of the invention is derived, flank the gene. The term therefore includes, for example, a recombinant DNA which is incorporated into a vector; into an autonomously replicating plasmid or virus; or into the genomic DNA of a prokaryote or eukaryote; or which exists as a separate molecule (e.g., a cDNA or a genomic or cDNA fragment produced by PCR or restriction endonuclease digestion) independent of other sequences. It also includes a recombinant DNA which is part of a hybrid gene encoding additional polypeptide sequence.

By "transformed cell" is meant a cell into which (or into an ancestor of which) has been introduced, by means of

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recombinant DNA techniques, a DNA molecule encoding (as used herein) an IAP polypeptide.

By "transgene" is meant any piece of DNA which is inserted by artifice into a cell, and becomes part of the genome of the organism which develops from that cell. Such a transgene may include a gene which is partly or entirely heterologous (i.e., foreign) to the transgenic organism, or may represent a gene homologous to an endogenous gene of the organism.

By "transgenic" is meant any cell which includes a DNA sequence which is inserted by artifice into a cell and becomes part of the genome of the organism which develops from that cell. As used herein, the transgenic organisms are generally transgenic mammalian (e.g., rodents such as rats or mice) and the DNA (transgene) is inserted by artifice into the nuclear genome.

By "transformation" is meant any method for introducing foreign molecules into a cell. Lipofection, calcium phosphate precipitation, retroviral deliver, electroporation and biolistic transformation are just a few of the teachings which may be used. For example, Biolistic transformation is a method for introducing foreign molecules into a cell using velocity driven microprojectiles such as tungsten or gold particles. Such velocity-driven methods originate from pressure bursts which include, but are not limited to, helium-driven, air-driven, and gunpowder-driven techniques. Biolistic transformation may be applied to the transformation or transfection of a wide variety of cell types and intact tissues including, without limitation, intracellular organelles (e.g., and mitochondria and chloroplasts), bacteria, yeast, fungi, algae, animal tissue, and cultured cells.

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By "positioned for expression" is meant that the DNA molecule is positioned adjacent to a DNA sequence which directs transcription and translation of the sequence (i.e., facilitates the production of, e.g., an IAP polypeptide, a recombinant protein or a RNA molecule).

By "reporter gene" is meant a gene whose expression may be assayed; such genes include, without limitation,  $\beta$ -glucuronidase (GUS), luciferase, chloramphenicol transacetylase (CAT), and  $\beta$ -galactosidase.

By "promoter" is meant minimal sequence sufficient to direct transcription. Also included in the invention are those promoter elements which are sufficient to render promoter-dependent gene expression controllable for cell-type specific, tissue-specific or inducible by external signals or agents; such elements may be located in the 5' or 3' regions of the native gene.

By "operably linked" is meant that a gene and a regulatory sequence(s) are connected in such a way as to permit gene expression when the appropriate molecules (e.g., transcriptional activator proteins) are bound to the regulatory sequence(s).

By "conserved region" is meant any stretch of six or more contiguous amino acids exhibiting at least 30%, preferably 50%, and most preferably 70% amino acid sequence identity between two or more of the IAP family members, (e.g., between human HIAP1, HIAP2, and XIAP). Examples of preferred conserved regions are shown (as boxed or designated sequences) in Figures 5-7 and Tables 1 and 2, and include, without limitation, BIR domains and ring zinc finger domains.

By "detectably-labelled" is meant any means for marking and identifying the presence of a molecule, e.g., an oligonucleotide probe or primer, a gene or fragment thereof,

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or a cDNA molecule. Methods for detectably-labelling a molecule are well known in the art and include, without limitation, radioactive labelling (e.g., with an isotope such as <sup>32</sup>P or <sup>35</sup>S) and nonradioactive labelling (e.g., chemiluminescent labelling, e.g., fluorescein labelling).

By "purified antibody" is meant antibody which is at least 60%, by weight, free from proteins and naturally-occurring organic molecules with which it is naturally associated. Preferably, the preparation is at least 75%, more preferably 90%, and most preferably at least 99%, by weight, antibody, e.g., an IAP specific antibody. A purified antibody may be obtained, for example, by affinity chromatography using recombinantly-produced protein or conserved motif peptides and standard techniques.

By "specifically binds" is meant an antibody which recognizes and binds a protein but which does not substantially recognize and bind other molecules in a sample, e.g., a biological sample, which naturally includes protein.

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

#### <u>Detailed Description</u>

The drawings will first be described.

## 25 <u>Drawings</u>

Fig. 1 is the human xiap cDNA sequence and the XIAP polypeptide sequence (SEQ ID NOS:3, 4).

Fig. 2 is the human hiap1 cDNA sequence and the HIAP1 polypeptide sequence (SEQ ID NOS:5, 6).

Fig. 3 is the human hiap2 cDNA sequence and the HIAP2 polypeptide sequence (SEQ ID NOS:7, 8). The sequence absent in the hiap2-G variant is boxed.

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Fig. 4 is the murine xiap cDNA sequence and encoded murine XIAP polypeptide sequence (SEQ ID NOS:9, 10).

Fig. 5 is the murine hiap1 cDNA sequence and the encoded murine HIAP1 polypeptide sequence (SEQ ID NOS:39, 40).

Fig. 6 is the murine hiap2 cDNA sequence and the encoded murine HIAP2 polypeptide SEQ ID NOS:41, 42).

Fig. 7 shows the alignment of the BIR domains of IAP proteins (SEQ ID NOS: 11 and 14-31).

Fig. 8 is the alignment of human IAP polypeptides with diap, cp-iap, and the consensus sequence (SEQ ID NOS:4, 6, 8, 10, 12, and 13).

Fig. 9 shows the alignment of the Ring Zinc Finger domains of IAP proteins (SEQ ID NOS: 32-38).

Fig. 10 is a Northern blot showing human hiap1 and hiap2 mRNA expression in human tissues.

Fig. 11 is a Northern blot showing human hiap2 mRNA expression in human tissues.

Fig. 12 is a Northern blot showing human xiap mRNA expression in human tissues.

Fig. 13A and 13B are agarose gels showing apoptic DNA ladders and RT PCR products using hiap1 and hiap2 specific probes in HIV infected T cells.

Fig. 14A - 14D are graphs showing apoptosis suppression by XIAP, HIAP1, HIAP2, bcl-2m, smn and 6-myc.

# I. IAP Polypeptides and Genes Encoding IAP polypeptides

We have discovered a new class of mammalian proteins which modulate apoptosis (IAPs) and the genes which encode these proteins. The IAP proteins are characterized by the presence of a  $\underline{r}$ ing  $\underline{z}$ inc  $\underline{f}$ inger (RZF) domain (Fig. 9) and at least one BIR domain as defined by the boxed consensus sequences in Figs. 7 and 8 and by the sequence domains

provided in Tables 1 and 2. As examples of the IAP proteins we provide the cDNA sequences and amino acid sequences for these new human and murine apoptosis inhibitors, HIAP1, HIAP2, and XIAP. Additional members of the mammalian IAP family (including homologs from other species and mutant sequences) may be isolated using standard cloning techniques and the conserved amino acid sequences, primers and probes provided herein and known in the art.

This application is related to U.S. Serial Number 08/511,485, filed August 4, 1995. U.S.S.N 08/511,485 is hereby incorporated by reference.

TABLE 1
NUCLEOTIDE POSITION OF CONSERVED DOMAINS\*

	BIR-1	BIR-2	BIR-3	Ring Zinc Finger
h-xiap	109 - 312	520 - 723	826 - 1023	1348-1485
m-xiap	202 - 405	613 - 816	916 - 1113	1438-1575
h-hiap1	273 - 476	693 - 893	951 - 1154	1824-1961
m-hiap1	251 - 453	670 - 870	928 - 1131	1795-1932
h-hiap2	373 - 576	787 - 987	1042-1245	1915-2052
m-hiap2	215 - 418	608 - 808	863 - 1066	1763-1876

Positions indicate correspond to those shown in Figs. 1-4.

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TABLE 2

AMINO ACID POSITION OF CONSERVED DOMAINS\*

	BIR-1	BIR-2	BIR-3	Ring Zinc Finger
h-Xiap	26 - 93	163 - 230	265 - 330	439 - 484
m-Xiap	26 - 93	163 - 230	264 - 329	438 - 483
h-Hiap1	29 - 96	169 - 235	255 - 322	546 - 591
m-Hiap1	29 - 96	169 - 235	255 - 322	544 - 589
h-Hiap2	46 - 113	184 - 250	269 - 336	560 - 605
m-Hiap2	25 - 92	156 - 222	241 - 308	541 - 578

Positions indicate correspond to those shown in Figs. 1-4.

Recognition of this mammalian IAP family has provided emergent patterns of protein structure. Recognition of these patterns has also allowed us assign the function of a modulator of apoptosis to a drosophila gene product of previously unknown function (Genbank Accession Number M96581). The amino acid sequence of this protein, termed diap, is shown in Fig. 8 for comparison.

The IAP proteins may be used to inhibit the apoptosis which occurs as part of disease or disorder processes. For example, IAP polypeptides or nucleic acid encoding IAP polypeptides may be administered for the treatment of or prevention of apoptosis which occurs as a part of AIDS, neurodegenerative diseases, ischemic injury, toxin-induced liver disease and myelodysplastic syndromes. Nucleic acid encoding the IAP polypeptide may also be provided to inhibit apoptosis.

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## II. Cloning of IAP Genes

# A. XIAP

Our search for human genes potentially involved in apoptosis has resulted in the identification of an x-linked sequence tag site (STS) in the GenBank which demonstrated strong homology with the conserved RZF domain of CpIAP and OpIAP, the two baculovirus genes known to inhibit apoptosis (Clem et al., Mol. Cell Biol., 14:5212-5222, (1994); and Birnbaum et al, J. Virol. 68:2521-8, (1994)). Screening a human fetal brain ZapII cDNA library (Stratagene, La Jolla, CA) with this STS resulted in the identification and cloning of xiap (for X-linked Inhibitor of apoptosis protein gene). The human gene has a 1.7 kb coding sequence that includes three BIR (baculovirus inhibitor of apoptosis repeat (Crook et al., J. Virol. 67:2168-74, (1993), Clem et al., Science 254:1388-90, (1991); and Birnbaum et al., J. Virol., 68:2521-8, (1994)) domains and a zinc finger. Northern analysis with xiap reveals a greater than 7kb message expressed in different tissues particularly liver and kidney (Fig. 12). The large size of the transcript reflects large 5' and 3' untranslated regions.

# B. HUMAN HIAP1 and HIAP2

The hiap1 and hiap2 genes were cloned by screening a human liver library (Stratagene) with a probe including the whole xiap coding region at low stringency (40°C wash, 2xssc, 10% SDS) (Figs. 2 and 3). hiap1 and hiap2 were also independently detected using a probe derived from a expressed sequence tag (EST) (GenBank Accession No. T96284) which includes a portion of a BIR domain. This EST was originally isolated by the PCR amplification of a cDNA library using the EST-specific primers. The derived probe

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was then used to screen the human liver cDNA library for full length hiap coding sequences. We have subsequently detected a third DNA which includes the hiap2 sequence which appears to lack one exon, presumably due to alternative mRNA splicing (see boxed region in Fig. 3). Figures 8 and 9 show hiap1 and hiap2 expression in human tissues as assayed by Northern Analysis.

### C. M-XIAP

Screening of a mouse embryo \(\lambda\)gt11 cDNA library (Clonetech, Palo Alto, CA) and a mouse FIX II genomic library with a \(xiap\) cDNA clones probe has resulted in the identification of 14 positive cDNA and two hybridizing genomic clones. A cDNA contig spanning 8.0 kb was constructed using 12 overlapping mouse clones. DNA sequencing revealed a coding sequence of about 1.7 kb. The mouse gene called \(m-xiap\) (for mouse x-linked inhibitor of apoptosis protein gene) shows striking amino acid homology with xiap at and around the initiation methionine, the stop codon, the three BIR domains and the zinc finger domain. As with the human gene, the mouse homologue contains large 5' and 3' UTRs predicted to result in a transcript as large as 7-8 kb.

Sequencing and restriction mapping of m-xiap can be used to further delineate the structure and genomic organization of m-xiap. Southern blot analysis and inverse PCR technique (Groden et al., Cell 66:589-600 (1991) can be employed to map exons and sequence exon-intron boundaries.

Antisera can be raised against a m-xiap fusion protein expressed in *Escherichia coli* using a bacterial expression system. The resulting antisera can be used along with Northern blot analysis to analyze the spatial and temporal expression of m-xiap in the mouse.

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# D. M-HIAP1 and M-HIAP2

The murine homologs to hiap1 and hiap2 were cloned and sequenced in the same general manner as m-xiap using the human hiap1 and hiap2 sequences as probes. Cloning of m-hiap1 and m-hiap2 provide further demonstrations of the case with which homologs from different species may be detected and obtained using the techniques provided herein and those generally known to one skilled in the art of molecular biology.

# 10 III. Cloning of Additional IAP Genes

Low stringency Southern blot hybridization of human genomic DNA using probes specific for xiap, hiap1 and hiap2 show bands which correspond to the other known human IAP sequences. In addition, these probes detect sequences which do not correspond to known IAP sequences. This result indicates that additional IAP sequences may be readily identified using low stringency hybridization. Examples of murine and human xiap, hiap1, and hiap2 specific primers which may be used to clone additional genes by RT PCR are shown in Table 5. Standard techniques including PCR and hybridization may be used to clone homologs and additional genes.

# IV. Characterization of IAP Apoptosis Modulating Activity

The apoptosis inhibiting capability of IAPs can be defined in an *in vitro* system know to detect alterations in apoptosis. Mammalian expression constructs carrying IAPs and their truncated forms can be introduced into various cell lines such as CHO, HIH 3T3, HL6O, Rat-1, or Jurkart cells, for example. In addition, SF21 insect cells may be used in which case the IAP gene is preferentially expressed using an insect heat shock promotor. Apoptosis will then be

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induced in transfected cells and controls employing standard methodologies (e.g. serum withdrawal and staurosporine). A survival index (ratio of surviving transfected cells to surviving control cells) will indicate the strength of each IAP construct in inhibiting apoptosis. These experiments can confirm the presence of apoptosis inhibiting or enhancing activity and, can help to determine the minimal functional region of an IAP. These methods may also be used in combination with compounds to identify compounds which modulate apoptosis via their effect on IAP expression.

Figs. 14A - 14D show specific examples of apoptosis suppression assays. Fig. 14A shows CHO survival following serum withdrawal. CHO cells were transfected via Lipofectace with 2  $\mu g$  of each of the following recombinant plasmids; pCDNA3-6myc-hiap-1, pCDNA3-6myc-hiap-2, pCDNA3-6myc-xiap, pCDNA3-6myc, pCDNA3-HA-smn, and pCDNA3-bcl-2. Oligonucleotide primers were synthesized to allow PCR amplification and cloning of the xiap, hiap-1 and hiap-2. Oligonucleotide primers were synthesized to allow PCR amplification and cloning of the xiap, hiap-1, and hiap-2 ORFs in pCDNA3 (Invitrogen). Each construct was modified to incorporate a synthetic myc tag encoding six repeats of the peptide sequence MEQKLISEEDL allowing detection of myc-IAP fusion proteins via monoclonal anti-myc antiserum (Egan, et al., Nature 363:45-51, 1993). Triplicate samples of cell lines in 24 well dishes were washed 5 times with serum free media and maintained in serum free conditions during the course of the experiment. Trypan blue exclusion counting of viable cells utilizing a hemocytometer was performed on samples at time zero, 24 hrs., 48 hrs., and 72 hrs., post serum withdrawal. Survival was calculated as a percentage of initial numbers. Numbers represent the average of three separate experiments performed in triplicate, +/- average

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deviation. Fig. 14B shows survival of CHO transfected cell lines following exposure to menadione. Cell lines were plated in 24 well dishes, allowed to grow overnight, then exposed for 1.5 hrs. to [20mM] menadione (Sigma).

Triplicate samples were harvested at the time of exposure and at 24 hrs. post exposure and assessed by trypan blue exclusion for survival. Data represents the average of three independent experiments, +/- average deviation. Fig. 14C shows survival of Rat-1 cells following staurosporine exposure. Rat-1 cells were transfected with the plasmids listed in a), with selection in [800 mg/ml] G418 media for two weeks. Cell lines were assessed for resistance to [1\mu M]staurosporine induced apoptosis for 5 hrs. Viable cell counts were obtained 24 hrs. post exposure via trypan blue exclusion counting of samples prepared in triplicate.

Numbers represent the average of two independent experiments, +/- average deviation. Fig. 14D shows Rat-1 cell lines were tested for resistance to [10 mM] menadione for 1.5 hrs., then counted at 18 hrs. post exposure.

Numbers represent the average of three experiments performed in triplicate, +/- average deviation.

Specific examples of apoptosis assays are also provided in the following references:

Lymphocyte: C.J. Li et al., "Induction of apoptosis in uninfected lymphocytes by HIV-1 Tat protein", Science, 268:429-431 (1995); D. Gibellini et al., "Tat-expressing Jurkat cells show an increased resistance to different apoptotic stimuli, including acute human immunodeficiency virus-type 1 (HIV-1) infection", Br. J. Haematol. 89:24-33, (1995); S.J. Martin et al., "HIV-1 infection of human CD4+ T cells in vitro. Differential induction of apoptosis in

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these cells." J. Immunol. 152:330-42, (1994); C. Terai et al., "Apoptosis as a mechanism of cell death in cultured T lymphoblasts acutely infected with HIV-1", J. Clin Invest., 87:1710-5, (1991); J. Dhein et al., "Autocrine T-cell suicide mediated by APO-1/(Fas/CD95)", Nature 373:438-441, (1995); P.D. Katsikis et al., "Fas antigen stimulation induces marked apoptosis of T lymphocytes in human immunodeficiency virus-infected individuals", J. Exp. Med. 1815:2029-2036, (1995); Michael O. Westendorp et al., Sensitization of T cells to CD95-mediated apoptosis by HIV-1 Tat and gp120", Nature, 375:497, (1995); DeRossi et al., Virology 198:234-44, (1994).

Fibroblasts: H. Vossbeck et al., "Direct transforming activity of TGF-beta on rat fibroblasts", Int. J. Cancer, 61:92-97, (1995); S. Goruppi et al., "Dissection of c-myc domains involved in S phase induction of NIH3T3 fibroblasts", Oncogene, 9:1537-44, (1994); A. Fernandez et al., "Differential sensitivity of normal and Ha-ras-transformed C3H mouse embryo fibroblasts to tumor necrosis factor: induction of bcl-2, c-myc, and manganese superoxide dismutase in resistant cells", Oncogene, 9:2009-17, (1994); E. A. Harrington et al., "c-Myc-induced apoptosis in fibroblasts in inhibited by specific cytokines", Embo J., 13:3286-3295, (1994); N. Itoh et al., "A novel protein domain required for apoptosis. Mutational analysis of human Fas antigen", J. Biol. Chem., 268:10932-7, (1993).

Neuronal Cells: G. Melino et al., "Tissue transglutaminase and apoptosis: sense and antisense transfection studies with human neuroblastoma cells", Mol. Cell. Biol., 14:6584-6596, (1994); D. M. Rosenbaum et al., "Evidence for hypoxia-induced, programmed cell death of

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cultured neurons", Ann. Neurol., 36:864-870, (1994); N. Sato et al., "Neuronal differentiation of PC12 cells as a result of prevention of cell death by bcl-2", J. Neurobiol, 25:1227-1234, (1994); G. Ferrari et al., "N-acetylcysteine (D- and L-stereoisomers) prevents apoptotic death of 5 neuronal cells", J. Neurosci., 1516:2857-2866, (1995); A. K. Talley et al., "Tumor necrosis factor alpha-induced apoptosis in human neuronal cells: protection by the antioxidant N-acetylcysteine and the genes bcl-2 and crmA", Mol. Cell Biol., 1585:2359-2366, (1995); A. K. Talley et 10 al., "Tumor Necrosis Factor Alpha-Induced Apoptosis in Human Neuronal Cells: Protection by the Antioxidant N-Acetylcysteine and the Genes bcl-2 and crmA", Mol. and Cell. Biol., 15:2359-2366, (1995); G. Walkinshaw et al., "Induction of apoptosis in catecholaminergic PC12 cells by 15 Implications for the treatment of Parkinson's disease.", J. Clin. Invest. 95:2458-2464, (1995).

Insect Cells: R. J. Clem et al., "Prevention of apoptosis by a baculovirus gene during infection of insect cells", Science, 254:1388-90, (1991); N. E. Crook et al., "An apoptosis-inhibiting baculovirus gene with a zinc finger-like motif", J. Virol., 67:2168-74, (1993); S. Rabizadeh et al., "Expression of the baculovirus p35 gene inhibits mammalian neural cell death", J. Neurochem., 61:2318-21, (1993); M. J. Birnbaum et al., "An apoptosis-inhibiting gene from a nuclear polyhidrosis virus encoding a polypeptide with Cys/His sequence motifs", J. Virol, 68:2521-8, (1994); R. J. Clem et al., "Control of programmed cell death by the baculovirus genes p35 and iap", Mol. Cell. Biol., 14:5212-5222, (1994).

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# V. Construction of a Transgenic Animal

Characterization of IAPs can provide information that allows for the development of an IAP knockout animal model, preferably mammal, most preferably a mouse, by homologous recombination. Similarly, an IAP overproducing animal may be produced by means of DNA sequence integration into the genome.

A replacement type targeting vector to create a knockout can be constructed using an isogenic genomic clone from a mouse strain, e.g. 129/Sv (Strategene LaJolla, CA). The targeting vector will be introduced into a J1 line of embryonic stem (ES) cells by electroporation to generate ES cell lines that carry a profoundly truncated form of an IAP. To generate chimeric founder mice, the targeted cell lines will be injected into a mouse blastula stage embryo. Heterozygote offspring will be interbred to homozygosity. Knockout mice may be constructed as a means of screening in vivo for therapeutic compounds which modulate apoptosis.

Animals having enhanced IAP expression may also be constructed using standard transgenic technologies.

# VI. IAP Protein Expression

IAP genes may be expressed in both prokaryotic and eukaryotic cell types. For those IAP's which increase apoptosis it may be desirable to express the protein under control of an inducible promotor for the purposes of protein production.

In general, IAP proteins according to the invention may be produced by transformation of a suitable host cell with all or part of a IAP-encoding cDNA fragment (e.g., the cDNA described above) in a suitable expression vehicle.

Those skilled in the field of molecular biology will understand that any of a wide variety of expression systems

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may be used to provide the recombinant protein. The precise host cell used is not critical to the invention. The IAP protein may be produced in a prokaryotic host (e.g., E. coli) or in a eukaryotic host (e.g., Saccharomyces cerevisiae, insect cells, e.g., Sf21 cells, or mammalian cells, e.g., COS 1, NIH 3T3, or HeLa cells). Such cells are available from a wide range of sources (e.g., the American Type Culture Collection, Rockland, MD; also, see, e.g., Ausubel et al., Current Protocols in Molecular Biology, John Wiley & Sons, New York, 1994). The method of transformation or transfection and the choice of expression vehicle will depend on the host system selected. Transformation and transfection methods are described, e.g., in Ausubel et al., (supra); expression vehicles may be chosen from those provided, e.g., in Cloning Vectors: A Laboratory Manual (P.H. Pouwels et al., 1985, Supp. 1987).

One preferred expression system is the baculovirus system (using, for example, the vector pBacPAK9) available from Clontech (Palo Alto, CA). If desired, this system may be used in conjunction with other protein expression techniques, for example, the myc tag approach described by Evan et al. (Mol. Cell Biol. 5:3610-3616, 1985).

Alternatively, a IAP protein is produced by a stably-transfected mammalian cell line. A number of vectors suitable for stable transfection of mammalian cells are available to the public, e.g., see Pouwels et al. (supra); methods for constructing such cell lines are also publicly available, e.g., in Ausubel et al. (supra). In one example, cDNA encoding the IAP protein is cloned into an expression vector which includes the dihydrofolate reductase (DHFR) gene. Integration of the plasmid and, therefore, the IAP protein-encoding gene into the host cell chromosome is selected for by inclusion of 0.01-300  $\mu\rm M$  methotrexate in the

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cell culture medium (as described in Ausubel et al., <u>supra</u>). This dominant selection can be accomplished in most cell types. Recombinant protein expression can be increased by DHFR-mediated amplification of the transfected gene.

Methods for selecting cell lines bearing gene amplifications are described in Ausubel et al. (<u>supra</u>); such methods generally involve extended culture in medium containing gradually increasing levels of methotrexate.

DHFR-containing expression vectors commonly used for this purpose include pCVSEII-DHFR and pAdD26SV(A) (described in Ausubel et al., <a href="mailto:supra">supra</a>). Any of the host cells described above or, preferably, a DHFR-deficient CHO cell line (e.g., CHO DHFR cells, ATCC Accession No. CRL 9096) are among the host cells preferred for DHFR selection of a stably-transfected cell line or DHFR-mediated gene amplification.

Once the recombinant IAP protein is expressed, it is isolated, e.g., using affinity chromatography. In one example, an anti-IAP protein antibody (e.g., produced as described herein) may be attached to a column and used to isolate the IAP protein. Lysis and fractionation of IAP protein-harboring cells prior to affinity chromatography may be performed by standard methods (see, e.g., Ausubel et al., supra).

Once isolated, the recombinant protein can, if desired, be further purified, e.g., by high performance liquid chromatography (see, e.g., Fisher, <u>Laboratory</u>

<u>Techniques In Biochemistry And Molecular Biology</u>, eds., Work and Burdon, Elsevier, 1980).

Polypeptides of the invention, particularly short

IAP protein fragments, can also be produced by chemical synthesis (e.g., by the methods described in <u>Solid Phase Peptide Synthesis</u>, 2nd ed., 1984 The Pierce Chemical Co., Rockford, IL).

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These general techniques of polypeptide expression and purification can also be used to produce and isolate useful IAP fragments or analogs (described herein).

#### VI. Anti-IAP Antibodies

To generate IAP-specific antibodies, a IAP coding sequence (i.e., amino acids 180-276) can be expressed as a C-terminal fusion with glutathione S-transferase (GST) (Smith et al., Gene 67:31-40, 1988). The fusion protein can be purified on glutathione-Sepharose beads, eluted with glutathione cleaved with thrombin (at the engineered cleavage site), and purified to the degree necessary for immunization of rabbits. Primary immunizations can be carried out with Freund's complete adjuvant and subsequent immunizations with Freund's incomplete adjuvant. Antibody titres are monitored by Western blot and immunoprecipitation analyses using the thrombin-cleaved IAP protein fragment of the GST-IAP fusion protein. Immune sera are affinity purified using CNBr-Sepharose-coupled IAP protein. Antiserum specificity is determined using a panel of unrelated GST proteins (including GSTp53, Rb, HPV-16 E6, and E6-AP) and GST-trypsin (which was generated by PCR using known sequences).

As an alternate or adjunct immunogen to GST fusion proteins, peptides corresponding to relatively unique hydrophilic regions of IAP may be generated and coupled to keyhole limpet hemocyanin (KLH) through an introduced C-terminal lysine. Antiserum to each of these peptides is similarly affinity purified on peptides conjugated to BSA, and specificity tested in ELISA and Western blots using peptide conjugates, and by Western blot and immunoprecipitation using IAP expressed as a GST fusion protein.

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Alternatively, monoclonal antibodies may be prepared using the IAP proteins described above and standard hybridoma technology (see, e.g., Kohler et al., Nature 256:495, 1975; Kohler et al., Eur. J. Immunol. 6:511, 1976; Kohler et al., Eur. J. Immunol. 6:292, 1976; Hammerling et al., In Monoclonal Antibodies and T Cell Hybridomas, Elsevier, NY, 1981; Ausubel et al., <a href="supra">supra</a>). Once produced, monoclonal antibodies are also tested for specific IAP recognition by Western blot or immunoprecipitation analysis (by the methods described in Ausubel et al., <a href="supra">supra</a>). Antibodies which specifically recognize IAP are considered to be useful in the invention; such antibodies may be used, e.g., in an immunoassay to monitor the level of IAP produced by a mammal (for example, to determine the amount or subcellular location of IAP).

Preferably, antibodies of the invention are produced using fragments of the IAP protein which lie outside highly conserved regions and appear likely to be antigenic, by criteria such as those provided by the Peptidestructure program of the Genetics Computer Group Sequence Analysis Package (Program Manual for the GCG Package, Version 7, 1991) using the algorithm of Jameson and Wolf (CABIOS 4:181 Specifically these regions, which are found between BIR1 and BIR2 of all the IAP proteins, are in hiap1 from amino acid 99 to 170, hiap2 from amino acid 123 to 184, xiap from 116 to 133 and m-xiap from 116 to 133. In one specific example, such fragments are generated by standard techniques of PCR and cloned into the pGEX expression vector (Ausubel et al., <u>supra</u>). Fusion proteins are expressed in <u>E. coli</u> and purified using a glutathione agarose affinity matrix as described in Ausubel et al. (supra). To attempt to minimize the potential problems of low affinity or specificity of antisera, two or three such fusions are generated for each

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protein, and each fusion is injected into at least two rabbits. Antisera are raised by injections in a series, preferably including at least three booster injections.

# VII. Identification of Molecules that Modulate IAP Protein Expression

Isolation of the IAP cDNAs also facilitates the identification of molecules which increase or decrease IAP expression. According to one approach, candidate molecules are added at varying concentrations to the culture medium of cells expressing IAP mRNA. IAP expression is then measured, for example, by standard Northern blot analysis (Ausubel et al., <a href="supra">supra</a>) using a IAP cDNA (or cDNA fragment) as a hybridization probe (see also Table 5). The level of IAP expression in the presence of the candidate molecule is compared to the level measured for the same cells in the same culture medium but in the absence of the candidate molecule.

If desired, the effect of candidate modulators on expression may, in the alternative, be measured at the level of IAP protein production using the same general approach and standard immunological detection techniques, such as Western blotting or immunoprecipitation with a IAP-specific antibody (for example, the IAP antibody described herein).

Candidate modulators may be purified (or substantially purified) molecules or may be one component of a mixture of compounds (e.g., an extract or supernatant obtained from cells; Ausubel et al., <a href="supra">supra</a>). In a mixed compound assay, IAP expression is tested against progressively smaller subsets of the candidate compound pool (e.g., produced by standard purification techniques, e.g., HPLC or FPLC) until a single compound or minimal compound mixture is demonstrated to modulate IAP expression.

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Alternatively, or in addition, candidate compounds may be screened for those which modulate IAP apoptosis inhibiting activity. In this approach, the degree of apoptosis in the presence of a candidate compound is compared to the degree of apoptosis in its absence, under equivalent conditions. Again, such a screen may begin with a pool of candidate compounds, from which one or more useful modulator compounds are isolated in a step-wise fashion. Apoptosis activity may be measured by any standard assay, for example, those described herein.

Another method for detecting compounds which modulate IAP polypeptide activity is to screen for compounds which physically interact with a given IAP polypeptide. Such compounds may be detected using adaptations of the interaction trap expression systems known in the art. Such systems detect protein interactions using a transcriptional activation assay and are generally described in Gyuris et al., Cell 75:791-803 (1993), and Field and Song, Nature 340:245-246, (1989), and are commercially available from Clonetech (Palo Alto, CA). In addition, PCT Publication WO 95/28497 (hereby incorporated by reference) describe a method for detecting proteins involved in apoptosis by virtue of their interaction with Bcl-2 using such an interaction trap assay. A similar method may be exploited to identify proteins and other compounds which interact with the IAP polypeptides.

Candidate IAP modulators include peptide as well as non-peptide molecules (e.g., peptide or non-peptide molecules found, e.g., in a cell extract, mammalian serum, or growth medium on which mammalian cells have been cultured).

A molecule which promotes an increase in IAP expression or IAP activity is considered particularly useful

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in the invention; such a molecule may be used, for example, as a therapeutic to increase cellular levels of IAP and thereby exploit the effect of IAP polypeptides for the inhibition of apoptosis.

A molecule which decreases IAP activity (e.g., by decreasing gene expression or polypeptide activity) may be useful for decreasing cell proliferation. Such uses include treatment of neoplasms (see Table 3, below) or other cell proliferative diseases.

Modulators found to be effective at the level of IAP expression or activity may be confirmed as useful in animal models and, if successful, may be used as anti-cancer therapeutics for either the inhibition or the enhancement of apoptosis, as appropriate.

## IX. IAP Therapy

Because expression levels of IAP genes correlates with the levels of apoptosis, the IAP gene also finds use in anti-apoptosis gene therapy. In particular, to sustain neuronal cells, lymphocytes (T-cells and B-cells), or cells exposed to ischemic injury, a functional IAP gene may be introduced into cells at the sites predicted to undergo undesirable apoptosis.

Retroviral vectors, adenoviral vectors, adenoassociated viral vectors, or other viral vectors with the
appropriate tropism for cells likely to be involved in
apoptosis (for example, epithelial cells) may be used as a
gene transfer delivery system for a therapeutic IAP gene
construct. Numerous vectors useful for this purpose are
generally known (Miller, Human Gene Therapy 15-14, 1990;
Friedman, Science 244:1275-1281, 1989; Eglitis and Anderson,
BioTechniques 6:608-614, 1988; Tolstoshev and Anderson,
Current Opinion in Biotechnology 1:55-61, 1990; Sharp, The

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Lancet 337:1277-1278, 1991; Cornetta et al., Nucleic Acid Research and Molecular Biology 36:311-322, 1987; Anderson, Science 226:401-409, 1984; Moen, Blood Cells 17:407-416, 1991; and Miller and Rosman, Biotechniques 7:980-990, 1989; Le Gal La Salle et al., Science 259:988-990, 1993; and Johnson, Chest 107:77S-83S, 1995). Retroviral vectors are particularly well developed and have been used in clinical settings (Rosenberg et al., N. Engl. J. Med 323:370, 1990; Anderson et al., U.S. Pat. No. 5,399,346).

Non-viral approaches may also be employed for the introduction of therapeutic DNA into cells otherwise predicted to undergo apoptosis. For example, IAP may be introduced into a neuronal cell or a T-cell by the techniques of lipofection (Felgner et al., Proc. Natl. Acad. Sci. USA 84:7413, 1987; Ono et al., Neuroscience Lett 117:259, 1990; Brigham et al., Am. J. Med. Sci. 298:278, 1989; Staubinger and Papahadjopoulos, Meth. Enz. 101:512, 1983); asialorosonucoid-polylysine conjugation (Wu and Wu, J. Biol. Chem. 263:14621, 1988; Wu et al., J. Biol. Chem. 264:16985, 1989); or, less preferably, microinjection under surgical conditions (Wolff et al., Science 247:1465, 1990).

For any of the above approaches, the therapeutic IAP DNA construct is preferably applied to the site of the predicted apoptosis event (for example, by injection), but may also be applied to tissue in the vicinity of the predicted apoptosis event or even to a blood vessel supplying the cells predicted to undergo apoptosis.

In the gene therapy constructs, IAP cDNA expression is directed from any suitable promoter (e.g., the human cytomegalovirus, simian virus 40, or metallothionein promoters), and its production is regulated by any desired mammalian regulatory element. For example, if desired, enhancers known to direct preferential gene expression in

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neural cells or T-cells may be used to direct IAP expression. Such enhancers include, without limitation, those enhancers which are characterized as tissue or cell specific in their expression.

Alternatively, if a IAP genomic clone is utilized as a therapeutic construct (for example, following its isolation by hybridization with the IAP cDNA described above), IAP expression is regulated by its cognate regulatory sequences or, if desired, by regulatory sequences derived from a heterologous source, e.g., any of the promoters or regulatory elements described above.

Less preferably, IAP gene therapy is accomplished by direct administration of the IAP mRNA to a cell predicted to undergo apoptosis. This mRNA may be produced and isolated by any standard technique, but is most readily produced by in vitro transcription using a IAP cDNA under the control of a high efficiency promoter (e.g., the T7 promoter). Administration of IAP mRNA to malignant cells is carried out by any of the methods for direct nucleic acid administration described above.

Ideally, the production of IAP protein by any gene therapy approach described above results in a cellular level of IAP that is at least equivalent to the normal, cellular level of IAP in an unaffected individual. Treatment by any IAP-mediated gene therapy approach may be combined with more traditional therapies.

Another therapeutic approach included within the invention involves direct administration of recombinant IAP protein, either to the site of a predicted apoptosis event (for example, by injection) or systemically by any conventional recombinant protein administration technique. The actual dosage of IAP depends on a number of factors, including the size and health of the individual patient,

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but, generally, between 0.1mg and 100mg inclusive are administered per day to an adult in any pharmaceutically-acceptable formulation.

# X. Administration of IAP polypeptides, IAP genes, or modulators of IAP synthesis or function

A IAP protein, gene, or modulator may be administered with a pharmaceutically-acceptable diluent, carrier, or excipient, in unit dosage form. Conventional pharmaceutical practice may be employed to provide suitable formulations or compositions to administer IAP to patients suffering from or presymptomatic for a IAP-associated carcinoma. Any appropriate route of administration may be employed, for example, parenteral, intravenous, subcutaneous, intramuscular, intracranial, intraorbital, ophthalmic, intraventricular, intracapsular, intraspinal, intracisternal, intraperitoneal, intranasal, aerosol, or oral administration. Therapeutic formulations may be in the form of liquid solutions or suspensions; for oral administration, formulations may be in the form of tablets or capsules; and for intranasal formulations, in the form of powders, nasal drops, or aerosols.

Methods well known in the art for making formulations are found in, for example, "Remington's Pharmaceutical Sciences." Formulations for parenteral administration may, for example, contain excipients, sterile water, or saline, polyalkylene glycols such as polyethylene glycol, oils of vegetable origin, or hydrogenated napthalenes. Biocompatible, biodegradable lactide polymer, lactide/glycolide copolymer, or polyoxyethylene-polyoxypropylene copolymers may be used to control the release of the compounds. Other potentially useful parenteral delivery systems for IAP modulatory compounds

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include ethylene-vinyl acetate copolymer particles, osmotic pumps, implantable infusion systems, and liposomes. Formulations for inhalation may contain excipients, for example, lactose, or may be aqueous solutions containing, for example, polyoxyethylene-9-lauryl ether, glycocholate and deoxycholate, or may be oily solutions for administration in the form of nasal drops, or as a gel.

If desired, treatment with a IAP protein, gene, or modulatory compound may be combined with more traditional therapies for the disease such as surgery, radiation, or chemotherapy for cancers; surgery, steroid therapy, and chemotherapy for autoimmune diseases; antiviral therapies for AIDS; and for example, TPA for ischemic injury.

## XI. Detection of A Condition Involving Altered Apoptosis

IAP polypeptides and nucleic acid sequences find diagnostic use in the detection or monitoring of conditions involving aberrant levels of apoptosis. For example, decrease expression of IAP may be correlated with enhanced apoptosis in humans (see XII, below). Accordingly, a decrease or increase in the level of IAP production may provide an indication of a deleterious condition. Levels of IAP expression may be assayed by any standard technique. For example, its expression in a biological sample (e.g., a biopsy) may be monitored by standard Northern blot analysis or may be aided by PCR (see, e.g., Ausubel et al., supra; PCR Technology: Principles and Applications for DNA Amplification, ed., H.A. Ehrlich, Stockton Press, NY; and Yap and McGee, Nucl. Acids. Res. 19:4294, 1991).

Alternatively, a patient sample may be analyzed for one or more mutations in the IAP sequences using a mismatch detection approach. Generally, these techniques involve PCR amplification of nucleic acid from the patient sample,

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followed by identification of the mutation (i.e., mismatch) by either altered hybridization, aberrant electrophoretic gel migration, binding or cleavage mediated by mismatch binding proteins, or direct nucleic acid sequencing. Any of these techniques may be used to facilitate mutant IAP detection, and each is well known in the art; examples of particular techniques are described, without limitation, in Orita et al., Proc. Natl. Acad. Sci. USA 86:2766-2770, (1989); and Sheffield et al., Proc. Natl. Acad. Sci. USA 86:232-236, (1989).

In yet another approach, immunoassays are used to detect or monitor IAP protein in a biological sample. specific polyclonal or monoclonal antibodies (produced as described above) may be used in any standard immunoassay format (e.g., ELISA, Western blot, or RIA assay) to measure IAP polypeptide levels; again comparison is to wild-type IAP levels, and a decrease in IAP production is indicative of a condition involving increased apoptosis. Examples of immunoassays are described, e.g., in Ausubel et al., supra. Immunohistochemical techniques may also be utilized for IAP detection. For example, a tissue sample may be obtained from a patient, and a section stained for the presence of IAP using an anti-IAP antibody and any standard detection system (e.g., one which includes a secondary antibody conjugated to horseradish peroxidase). General guidance regarding such techniques can be found in, e.g., Bancroft and Stevens (Theory and Practice of Histological Techniques, Churchill Livingstone, 1982) and Ausubel et al. (supra).

In one preferred example, a combined diagnostic

method may be employed that begins with an evaluation of IAP protein production (for example, by immunological techniques or the protein truncation test (Hogerrorst, F.B.L., et al., Nature Genetics 10:208-212 (1995) and also includes a

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nucleic acid-based detection technique designed to identify more subtle IAP mutations (for example, point mutations). As described above, a number of mismatch detection assays are available to those skilled in the art, and any preferred technique may be used (see above). By this approach, mutations in IAP may be detected that either result in loss of IAP expression or loss of IAP biological activity. In a variation of this combined diagnostic method, IAP biological activity is measured as protease activity using any appropriate protease assay system (for example, those described above).

Mismatch detection assays also provide the opportunity to diagnose a IAP-mediated predisposition to diseases of apoptosis. For example, a patient heterozygous for an IAP mutation may show no clinical symptoms and yet possess a higher than normal probability of developing one or more types of neurodegenerative, myelodysplastic or ischemic diseases. Given this diagnosis, a patient may take precautions to minimize their exposure to adverse environmental factors (for example, UV exposure or chemical mutagens) and to carefully monitor their medical condition (for example, through frequent physical examinations). This type of IAP diagnostic approach may also be used to detect IAP mutations in prenatal screens.

The IAP diagnostic assays described above may be carried out using any biological sample (for example, any biopsy sample or bodily fluid or tissue) in which IAP is normally expressed (for example, the inhibition of apoptosis). Identification of a mutant IAP gene may also be assayed using these sources for test samples.

Alternatively, a IAP mutation, particularly as part of a diagnosis for predisposition to IAP-associated degenerative disease, may be tested using a DNA sample from any cell, for

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example, by mismatch detection techniques; preferably, the DNA sample is subjected to PCR amplification prior to analysis.

To demonstrate the utility of IAP gene sequences as diagnostics and prognostics for cancer we probed the Clonetech (La Jolla) Human Cancer Cell Line Multiple Tissue Northern Blot (#7757-1). As Table 3 shows, all cancer lines tested showed increased IAP expression relative to samples from non-cancerous control cell lines. xiap expression was particularly high in HeLa (S-3), chronic myelogenous leukemia (K-562), colorectal adenocarcinoma (SW-480) and melanoma (G-361) lines. hiap1 expression was extremely high in Burkitt's lymphoma and was also elevated in colorectal adenocarcinoma. hiap2 expression was particularly high in chronic myelogenous leukemia (K-562) and colorectal adenocarcinoma (SW-480).

In addition, we note that we have mapped hiap1 and hiap2 to human chromosome 11g23. This is a known hotspot for cancer causing mutations.

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TABLE 3
Northern Blot IAP RNA levels in Cancer Cells\*

	xiap	hiap1	hiap2
Promylocytic Leukemia HL-60	+	+	+
Hela S-3	+	+	+
Chronic Myclogenous Leukemia K-562	+++	+	+++
Lymphoblastic Leukemia MDLT-	+++	+	+
Burkitt's Lymphoma Raji	+	+(x10)	+
Colorectal Adenocarcinoma SW-480	+++	+++	+++
Lung Carcinoma A-549	+	+	+
Melanoma G-361	+++	+	+

\*Levels are indicated by a (+) and are the approximate increase in RNA levels relative to Northern blots of RNA from non-cancerous control cell lines. A single plus indicates an estimated increase of at least 1-fold

## XII. Treatment of HIV Infected Individuals

We have found that hiap1 and hiap 2 expression is
decreased significantly in HIV infected human cells. This
decrease precedes apoptosis. The result indicates that
administration of HIAP1, HIAP2, genes encoding these
proteins, or compounds which upregulate these genes can be
used to prevent T-cell attrition in HIV infected patients.

25 The following assay may also be used to screen for compounds

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which alter hiap1 and hiap2 expression and which also prevent apoptosis.

The experiments were preformed as follows: Cultured mature lymphocyte CD-4<sup>+</sup> T-cell lines (H9 labelled "a"; CEM/CM-3 labelled "b"; 6T-CEM labelled "c"; and Jurkat labelled "d" in Figs. 13A and 13B) were examined for apoptosis (Fig. 13A) and hiap gene expression (Fig. 13B). Control conditions are labelled as lane 1 in Fig. 13A and Fig. 13B. Lane 2 shows the result 24 hours after PHA/PMH (phytohemagglutinin, phorbol ester) mitogen stimulation. Lane 3 shows the result 24 hours after HIV strain III<sub>B</sub> infection. The "M" refers to standard DNA markers, the 123 bp ladder (Gibco-BRL) in Fig. 13B, and lambda HindIII ladder (Gibco-BRL) in Fig. A.

In Fig. 13A is a picture of ethidium bromide stained gel showing the presence of DNA ladders (as assayed by the test of Prigent et al., J. of Immun. Methods, 160:139-140, (1993), indicative of apoptosis. The sensitivity and degree of apoptosis of the four T-cell lines varies following mitogen stimulation and HIV infection.

For the experiment examining hiap gene expression, total RNA was prepared from the cultured cells and subject to a reverse transcriptase reaction using oligo-dT priming. The RT cDNA products were PCR amplified using specific primers (as shown in Table 5) for the detection of hiap2a, hiap2b and hiap 1. PCR conditions were routine (94°C melting for 1 minute, 55°C annealing for 2 minutes and 72°C extension for 1.5 minutes for 35 cycles) using a Perkin-Elmer 480 thermocycler. The Fig. 13B shows a picture of the RT-PCR products run on a 1% agarose gel stained with ethidium bromide. Absence of hiap2 transcripts is noted in all four cell lines 24 hours after HIV infection. In three of four cell lines (all except H9), the hiap1 gene is also

dramatically down-regulated after HIV infection. PHA/PMA mitogen stimulation also appears to decrease hiap gene expression, particularly for hiap2 and to a lesser extent, for hiap1.

The data from these experiments is summarized in the accompanying Table 5. The \(\beta\)-action gene expression was consistent in all cell lines tested, indicating that a flow in the RT-PCR assay does not account for the decreases in hiap gene expression.

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IAP Gene	Forward Primer (nucleotide position*)	Reverse Primer (nucleotide position*)	Size of Product (bp)
h-xiap	p2415 (876-896)	p2449 (1291-1311)	435
m-xiap	p2566 (458-478)	p2490 (994-1013)	555
h-hiapl	p2465 (827-847)	p2464 (1008-1038)	211
m-hiap1	p2687 (747-767)	p2684 (1177-1197)	450
hıap2	p2595 (1562-1585)	p2578 (2339-2363)	801 <sup>a</sup> 618 <sup>b</sup>
m-hiap2	p2693 (1751-1772)	p2734 (2078-2100)	349

\* Nucleotide position as determined from Figs. 1-4 for each IAP gene

<sup>&</sup>lt;sup>a</sup> PCR product size of hiap2a

b PCR product size of hiap2b

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Table 5

Apoptosis and hiap gene expression in cultured T-cells following mitogen stimulation or HIV infection.

Cell Line	Condition	Apoptosis	hiap1	hiap2
Н9	not stimulated PHA/PMA stimulated HIV infected	- + + + + +	+ + +	+/- +/- -
CEM/CM-3	not stimulated PHA/PMA stimulated HIV infected	- +/- +/-	+ + -	+/- - -
6T-CEM	not stimulated PHA/PMA stimulated HIV infected	- + /- +	+ -	+
Jurkat	not stimulated PHA/PMA stimulated HIV infected	- + +/-	+ + -	++ + -

#### XIII. Preventive Anti-Apoptotic Therapy

In a patient diagnosed to be heterozygous for an IAP mutation or to be susceptible to IAP mutations (even if those mutations do not yet result in alteration or loss of IAP biological activity), or a patent diagnosed as HIV positive, any of the above therapies may be administered before the occurrence of the disease phenotype. For example, the therapies may be provided to a patient who is HIV positive but does not yet show a diminished T-cell count or other signs of full-blown AIDS. In particular, compounds shown to increase IAP expression or IAP biological activity may be administered by any standard dosage and route of administration (see above). Alternatively, gene therapy using an IAP expression construct may be undertaken to

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reverse or prevent the cell defect prior to the development of the degenerative disease.

The methods of the instant invention may be used to reduce or diagnose the disorders described herein in any mammal, for example, humans, domestic pets, or livestock. Where a non-human mammal is treated or diagnosed, the IAP polypeptide, nucleic acid, or antibody employed is preferably specific for that species.

#### Other Embodiments

In other embodiments, the invention includes any protein which is substantially identical to a mammalian IAP polypeptides (Figs. 1-6; SEQ ID NO:1-42); such homologs include other substantially pure naturally-occurring mammalian IAP proteins as well as allelic variants; natural mutants; induced mutants; DNA sequences which encode proteins and also hybridize to the IAP DNA sequences of Figs. 1-6 (SEQ ID NOS:1-42) under high stringency conditions or, less preferably, under low stringency conditions (e.g., washing at 2X SSC at 40°C with a probe length of at least 40 nucleotides); and proteins specifically bound by antisera directed to a IAP polypeptide. The term also includes chimeric polypeptides that include a IAP portion.

The invention further includes analogs of any naturally-occurring IAP polypeptide. Analogs can differ from the naturally-occurring IAP protein by amino acid sequence differences, by post-translational modifications, or by both. Analogs of the invention will generally exhibit at least 85%, more preferably 90%, and most preferably 95% or even 99% identity with all or part of a naturally-occurring IAP amino acid sequence. The length of sequence comparison is at least 15 amino acid residues, preferably at least 25 amino acid residues, and more preferably more than

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35 amino acid residues. Modifications include in vivo and in vitro chemical derivatization of polypeptides, e.g., acetylation, carboxylation, phosphorylation, or glycosylation; such modifications may occur during polypeptide synthesis or processing or following treatment with isolated modifying enzymes. Analogs can also differ from the naturally-occurring IAP polypeptide by alterations in primary sequence. These include genetic variants, both natural and induced (for example, resulting from random mutagenesis by irradiation or exposure to ethanemethylsulfate or by site-specific mutagenesis as described in Sambrook, Fritsch and Maniatis, Molecular Cloning: A Laboratory Manual (2d ed.), CSH Press, 1989, or Ausubel et al., supra). Also included are cyclized peptides, molecules, and analogs which contain residues other than L-amino acids, e.g., D-amino acids or nonnaturally occurring or synthetic amino acids, e.g., B or y amino acids.

In addition to full-length polypeptides, the invention also includes IAP polypeptide fragments. As used herein, the term "fragment," means at least 20 contiguous amino acids, preferably at least 30 contiguous amino acids, more preferably at least 50 contiguous amino acids, and most preferably at least 60 to 80 or more contiguous amino acids. Fragments of IAP polypeptides can be generated by methods known to those skilled in the art or may result from normal protein processing (e.g., removal of amino acids from the nascent polypeptide that are not required for biological activity or removal of amino acids by alternative mRNA splicing or alternative protein processing events).

Preferable fragments or analogs according to the invention are those which facilitate specific detection of a IAP nucleic acid or amino acid sequence in a sample to be

diagnosed. Particularly useful IAP fragments for this purpose include, without limitation, the amino acid fragments shown in Table 2.

All publications and patent applications mentioned in this specification are herein incorporated by reference to the same extent as if each independent publication or patent application was specifically and individually indicated to be incorporated by reference.

What is claimed is:

#### Claims

- 1. Substantially pure nucleic acid encoding an IAP
- 2 polypeptide.
- 1 2. The nucleic acid of claim 1, wherein said
- 2 polypeptide comprises a ring zinc finger domain and at least
- 3 one BIR domain.
- 1 3. The nucleic acid of claim 2, wherein said
- 2 polypeptide has at least two BIR domains.
- 1 4. The nucleic acid of claim 3, wherein said
- 2 polypeptide has at least three BIR domains.
- 1 5. The nucleic acid of claim 1, wherein said DNA
- 2 contains the xiap gene.
- 1 6. The nucleic acid of claim 1, wherein said DNA
- 2 contains the hiap2 gene.
- 7. The nucleic acid of claim 1, wherein said DNA
- 2 contains the hiap1 gene.
- 1 8. The nucleic acid of claim 1, wherein said
- 2 nucleic acid is genomic DNA.
- 1 9. The nucleic acid of claim 1, wherein said
- 2 nucleic acid is cDNA.
- 1 10. The nucleic acid of claim 1, wherein said
- 2 nucleic acid is mammalian DNA.

- 1 11. The nucleic acid of claim 10, wherein said
- 2 mammalian DNA is human DNA.
- 1 12. The nucleic acid of claim 10, wherein said
- 2 mammalian DNA is murine DNA.
- 1 13. Substantially pure DNA having the sequence of
- 2 Fig. 5, or degenerate variants thereof, and encoding the
- 3 amino acid sequence of Fig. 5.
- 1 14. Substantially pure DNA having the sequence of
- 2 Fig. 6, or degenerate variants thereof, and encoding the
- 3 amino acid sequence of Fig. 6.
- 1 15. Substantially pure DNA having about 50% or
- 2 greater nucleotide sequence identity to the DNA sequence of
- 3 Fig. 5.
- 1 16. Substantially pure DNA having about 50% or
- 2 greater nucleotide sequence identity to the DNA sequence of
- 3 Fig. 6.
- 1 17. A purified DNA sequence substantially identical
- 2 to the DNA sequence shown in Fig. 5.
- 1 18. A purified DNA sequence substantially identical
- 2 to the DNA sequence shown in Fig. 6.
- 1 19. A substantially pure mammalian IAP polypeptide.
- 1 20. The polypeptide of claim 19, wherein said
- 2 polypeptide is the murine HIAP1 polypeptide.

- 1 21. The polypeptide of claim 19, wherein said
- 2 polypeptide is the murine HIAP2 polypeptide.
- 1 22. The polypeptide of claim 19, comprising an
- 2 amino acid sequence substantially identical to an amino acid
- 3 sequence shown in Fig. 5.
- 1 23. The polypeptide of claim 19, comprising an
- 2 amino acid sequence substantially identical to an amino acid
- 3 sequence shown in Fig. 6.
- 1 24. A therapeutic composition comprising as an
- 2 active ingredient an IAP polypeptide according to claim 19,
- 3 said active ingredient being formulated in a physiologically
- 4 acceptable carrier.
- 1 25. A method of inhibiting apoptosis in a mammal,
- 2 said method comprising:
- providing a cell of said mammal with a transgene
- 4 encoding an IAP polypeptide, said DNA positioned for
- 5 expression in said cell.
- 1 26. The method of claim 25 wherein said polypeptide
- 2 is murine HIAP1.
- 1 27. The method of claim 25 wherein said polypeptide
- 2 is murine HIAP2.
- 1 28. A method of detecting an IAP gene in an animal
- 2 cell, said method comprising:
- 3 contacting the DNA of claim 13 or a portion thereof
- 4 greater than about 18 nucleic acids in length with a
- 5 preparation of genomic DNA from said animal cell under

- 6 hybridization conditions providing detection of DNA
- 7 sequences having about 50% or greater nucleotide sequence
- 8 identity to the sequence of Fig. 5.
- 1 29. A method of detecting an IAP gene in an animal
- 2 cell, said method comprising:
- 3 contacting the DNA of claim 14 or a portion thereof
- 4 greater than about 18 nucleic acids in length with a
- 5 preparation of genomic DNA from said animal cell under
- 6 hybridization conditions providing detection of DNA
- 7 sequences having about 50% or greater nucleotide sequence
- 8 identity to the sequence of Fig. 6.
- 1 30. A method of producing an IAP polypeptide
- 2 comprising:
- providing a cell transformed with DNA encoding an
- 4 IAP polypeptide positioned for expression in said cell;
- 5 culturing said transformed cell under conditions for
- 6 expressing said DNA; and
- 7 isolating said IAP polypeptide.
- 1 31. The method of claim 30, wherein said IAP
- 2 polypeptide is murine HIAP1.
- 1 32. The method of claim 30, wherein said IAP
- 2 polypeptide is murine HIAP2.
- 1 33. A method of identifying a compound which
- 2 modulates apoptosis, said method comprising (a) providing a
- 3 cell expressing an IAP polypeptide; and (b) contracting said
- 4 cell with a candidate compound and monitoring the expression
- of an IAP gene, an alteration in the level of expression of

- 6 said gene indicating the presence of a compound which
- 7 modulates apoptosis.
- 1 34. The method of claim 33, wherein said IAP gene
- 2 is murine HIAP1.
- 1 35. The method of claim 33, wherein said IAP gene
- 2 is murine HIAP2.
- 1 36. A method for detecting a protein that interacts
- 2 with an IAP polypeptide comprising the steps of:
- 3 a. contacting under suitable conditions an IAP
- 4 protein with a compound suspected to be a modulator of
- 5 apoptosis; and
- 6 b. detecting the interaction of said compound with
- 7 said IAP polypeptide, wherein said interaction indicates
- 8 that said compound is involved in the modulation of
- 9 apoptosis.
- 1 37. The method of claim 36, wherein said IAP
- 2 polypeptide is HIAP1.
- 1 38. The method of claim 36, wherein said IAP
- 2 polypeptide is HIAP2.
- 1 39. The method of claim 36, wherein said IAP
- 2 polypeptide is XIAP.
- 1 40. The method of claim 36, wherein said
- 2 interaction is detected by measuring the transcriptional
- 3 activity of a reporter gene.

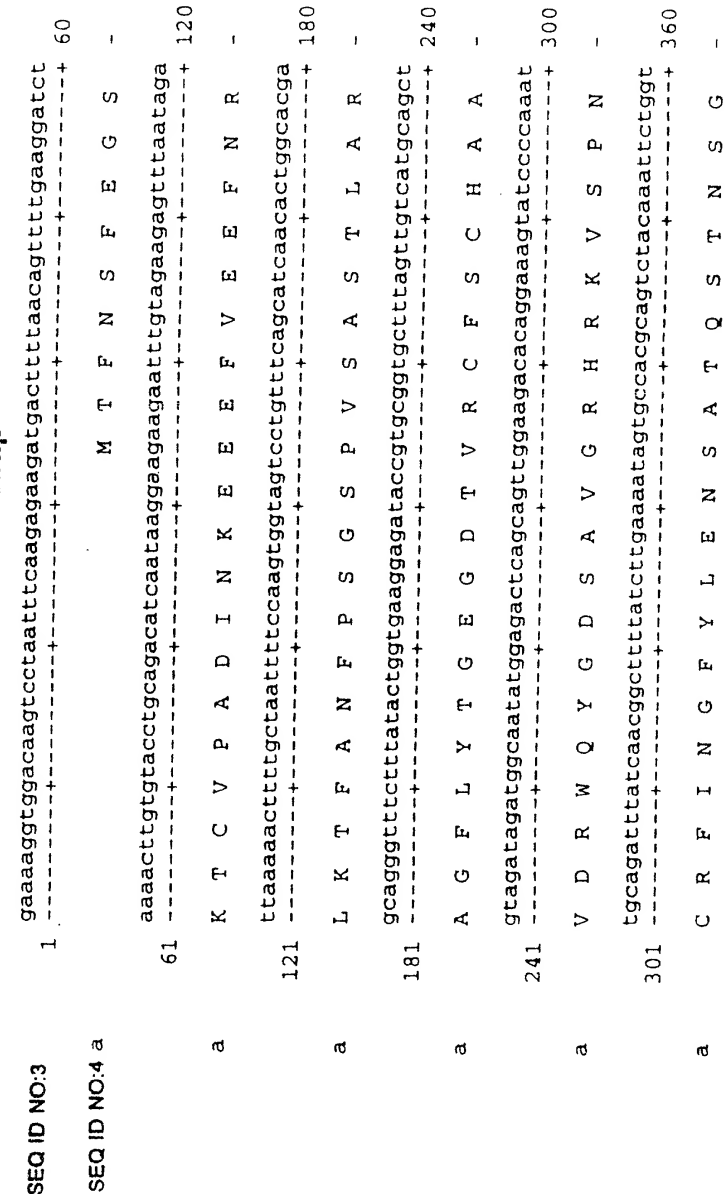
- 1 41. The method of claim 36, wherein said
- 2 interaction occurs in a yeast cell.
- 1 42. The method of claim 36, wherein said compound
- 2 is a polypeptide.
- 1 43. The method of claim 42, wherein said
- 2 polypeptide is expressed from a recombinant nucleic acid.
- 1 44. A method of diagnosing an increased liklihood
- 2 of a cell proliferative disease in a patient, said method
- 3 comprising detecting the level of IAP gene expression in
- 4 said patient.
- 1 45. A method of diagnosing an increased liklihood
- 2 of a cell proliferative disease in a patient, said method
- 3 comprising detecting the level of IAP polypeptide activity
- 4 in said patient.
- T 46. A transgenic rodent having a knockout mutation
- 2 in an IAP gene.
- 1 47. A transgenic rodent, said rodent having
- 2 additional copies of IAP nucleic acids added to its genome.

#### MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS

#### ABSTRACT OF THE DISCLOSURE

Disclosed is substantially pure DNA encoding mammalian IAP polypeptides; substantially pure polypeptides; and methods of using such DNA to express the IAP polypeptides in cells and animals to inhibit apoptosis. Also disclosed are conserved regions characteristic of the IAP family and primers and probes for the identification and isolation of additional IAP genes. In addition, methods for treating diseases and disorders involving apoptosis are provided.

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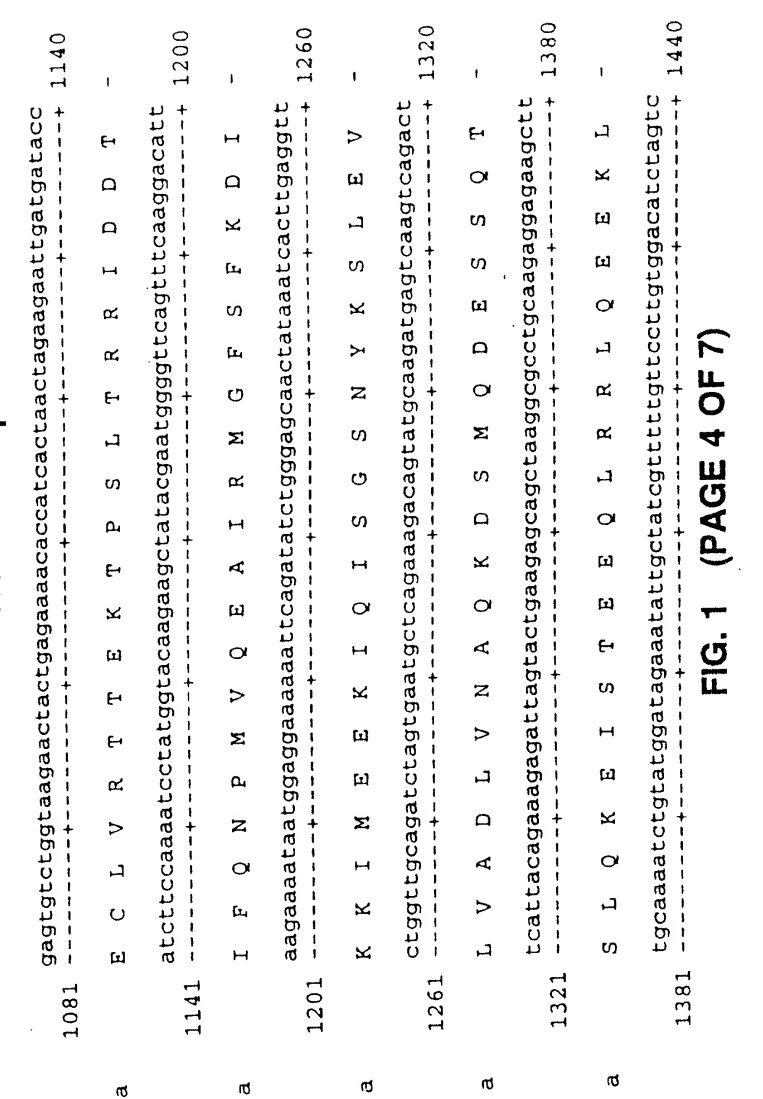
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atccagaatggtcagtacaagttgaaaactatctgggaagcagagatcattttgcctta 361++++++	A I Q N G Q Y K V E N Y L G S R D H F A L	gacaggccatctgagacacatgcagactatcttttgagaactgggcaggttgtagatata 421+++++++	a DRPSETHADYLLRTGQVVDI.	tcagacaccatatacccgaggaaccctgccatgtattgtgaagaagctagattaaagtcc 481+++++++	A SDTIYPAMYCEEARLKS.	tttcagaactggccagactatgctcacctaaccccaagagagttagcaagtgctggactc	A FONWPDYAHLTPRELASAGL	tactacacaggtattggtgaccaagtgcagtgcttttgttgtggtggaaaactgaaaat 601	A Y Y T G I G D Q V Q C F C C G G K L K N	tgggaaccttgtgatcgtgcctggtcagaacacaggcgacactttcctaattgcttcttt	A WEPCDRAWSEHRRHFPNCFF
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FIG. 1 (PAGE 2 OF 7)

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gttttgggccggaatcttaatactcgaagtgaatctgatgctgtgagttctga <b>ta</b> ggaat	VLGRNLNIRSESDAVSSDRN-	ttcccaaattcaacaaatcttccaagaaatccatccatggcagattatgaagcacggatc	FPNSTNLPRNPSMADYEARI-	tttacttttgggacatggatatactcagttaacaaggagcagcttgcaagagctggattt 841+++++++	FTFGTWIYSVNKEQLARAGF-	tatgetttaggtgaaggtgataaagtaaagtgettteaetgtggaggaggggetaaetgat 901	YALGEGDKVKCFHCGGGLTD -	tggaagcccagtgaagacccttgggaacaacatgctaaatggtatccagggtgcaaatat	WKPSEDPWEQHAKWYPGCKY	ctgttagaacagaagggacaagaatatataaacaatattcatttaactcattcacttgag 1021+++++++	LLEQKGOEYINNIHLTHSLE
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FIG. 1 (PAGE 3 OF 7)



	CKICMDRNIAIVFVPCGHLV -	
1441	acttgtaaacaatgtgctgaagcagttgacaagtgtcccatgtgctacacagtcattact	0
	TCKQCAEAVDKCPMCYTVIT -	
1501	ttcaagcaaaaattttttatgtcttaatctaactctatagtaggcatgttatgttgttct 	0
	FKOKIFMS*	
1561	tattaccctgattgaatgtgtgatgtgaactgactttaagtaatcaggattgaattccat ++++	0
1621	tagcatttgctaccaagtaggaaaaaatgtacatggcagtgttttagttggcaatata 	0 0
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2100	tgtaaagnnataaacacgnacntgtgcgaaatatntttgtaaagtgatttgccattnttg 2041+++++++	
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2040	gaaagatagagattgtttttagaggttggttgttgtgttttaggattctgtccattttct 1981	
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1980	atctccccaatcacatttgttttgtgtgaaaaaggaataaattgttccatgctggtg 1921	
1		ಹ
1920	tcttttcagataggcttaacaaatggagctttctgtatataaatgtggagattagagtta 1861	
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1860	attcatagtalactgatttctaagtgtaagtgaattaatcatctggattttttat 1801	

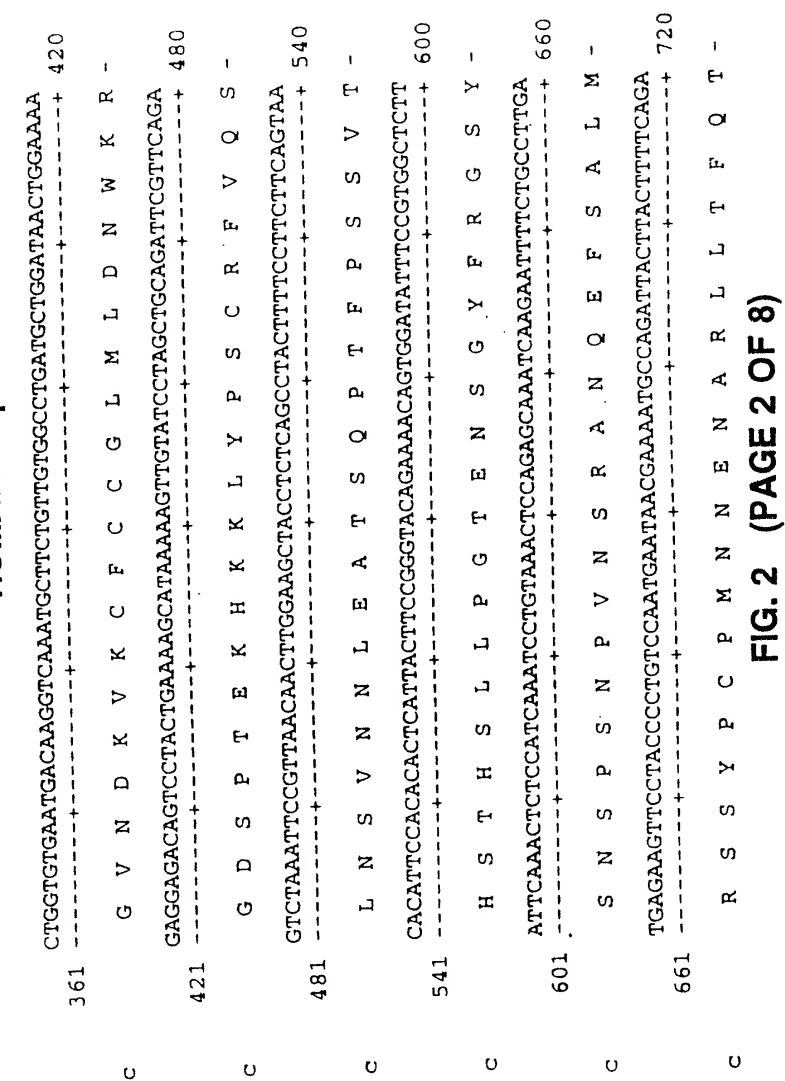
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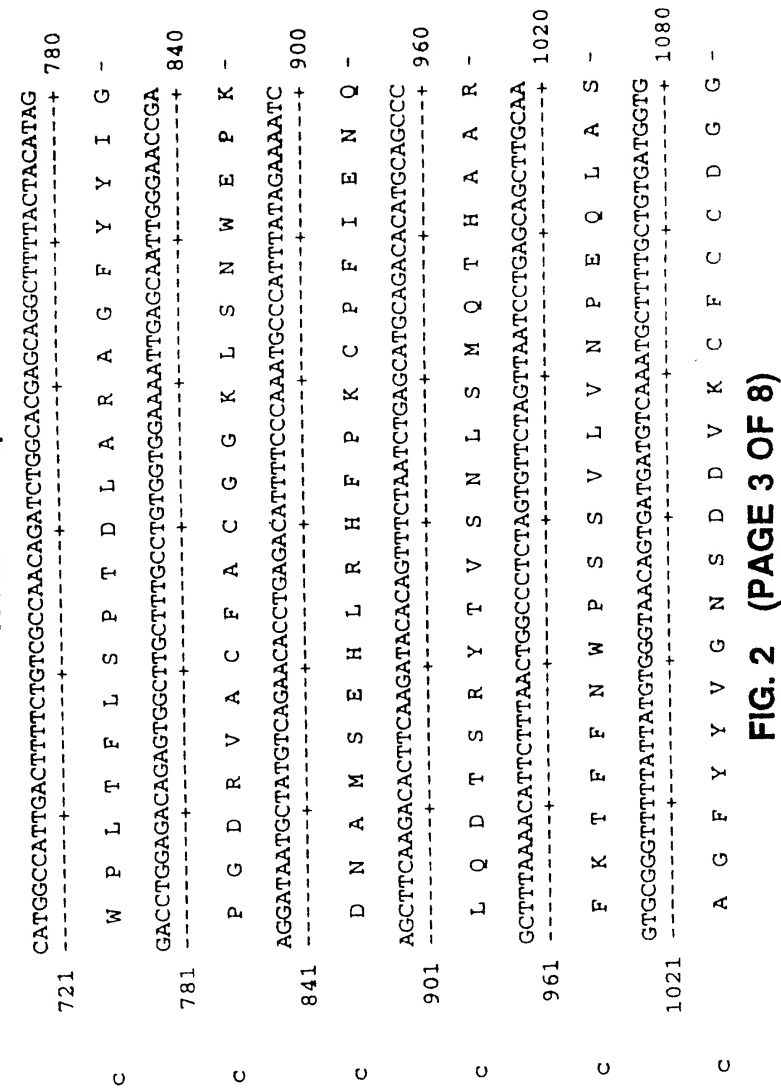
# HUMAN xiap

	ctgagtgctggggcactttn FIG. 1 (PAGE 7 OF 7)	2521	
į		М	ಥ
2520	gtnctcttcctagggagctgtnttgtttcccacccacccttccctt	2461	
ı		<b></b>	ಹ
2460	gtanaccccnaagggttttatggnaactaacatcagtaacctaacc	2401	
t			め
2400	naggggccttttcactttcnacttttttcattttgttctgttc	2341	
ı			d
2340	ttaaatgtggtttctcttcggggggggggggggggtttggggggg	2281	
1			ರ
2280	aagtatgtatgttnttaatatgcatagaacnanagatttggaaagatatacaccaaactg  ++++	2221	
1			ಹ
2220	nagatatgttaagtgtaaaatgcaagtggcnnnacactatgtatagtctgagccagatca	2161	

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SEQ ID NO:5	TCCTTGAGATGTATCAGTATTAGGATCTCCATGTTGGAACTCTAAATGCATAGA	09
O	AATGGAAATAATGGAAATTTTCATTTTGGCTTTTTCAGCCTAGTATTAAAACTGATAAAA	120
O	GCAAAGCCATGCACAAACTACCTCCCTAGAGAAAGGCTAGTCCCTTTTCTTCCCCATTC	180
		i
	ATTTCATTATGAACATAGTAGAAAACAGCATATTCTTATCAAATTTGATGAAAAGCGCCA	240
SEQ ID NO:6 C	MNIVENSIFLSNLMKSAN	1
	ACACGTTTGAAATACGACTTGTCATGTGAACTGTACCGAATGTCTACGTATTCCA	300
υ	TFELKYDLSCELYRMSTYST	l C.
	CTTTTCCTGCTGGGTTCCTGTCTCAGAAAGGAGTCTTGCTCGTGCTGGTTTCTATTACA	1 + 360
U	FPAGVPVSERSLARAGFYYT FIG. 2 (PAGE 1 OF 8)	i 5-4





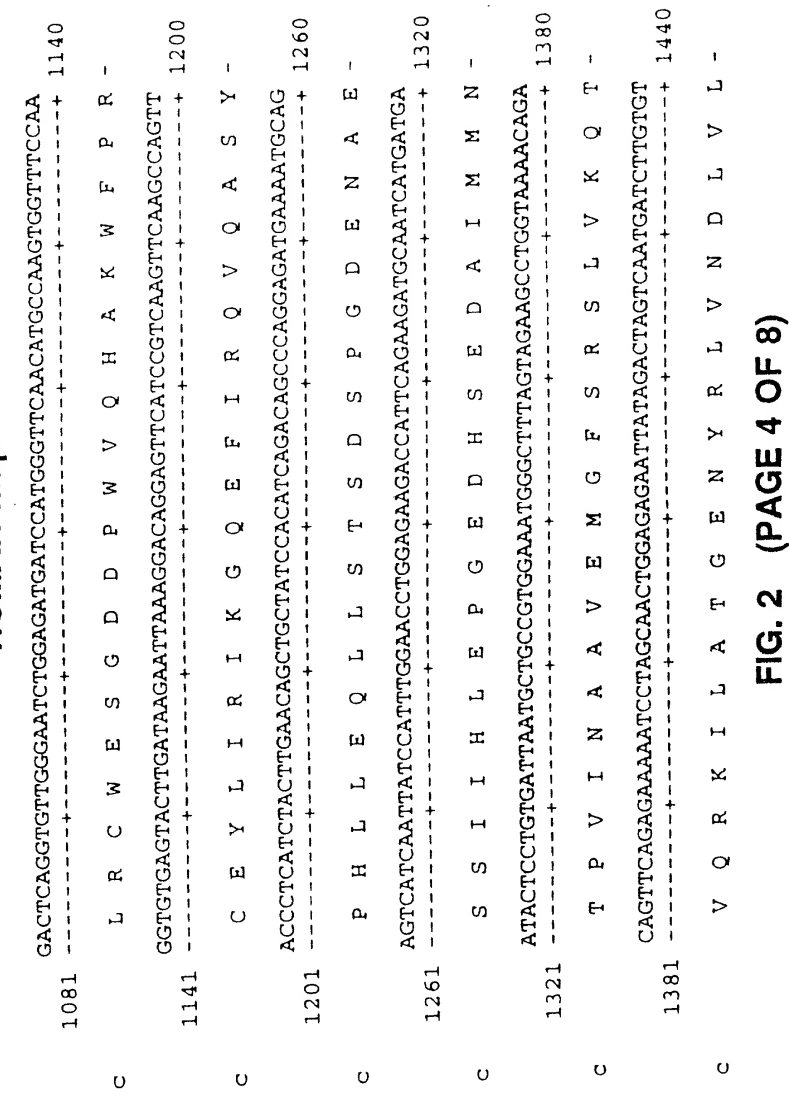
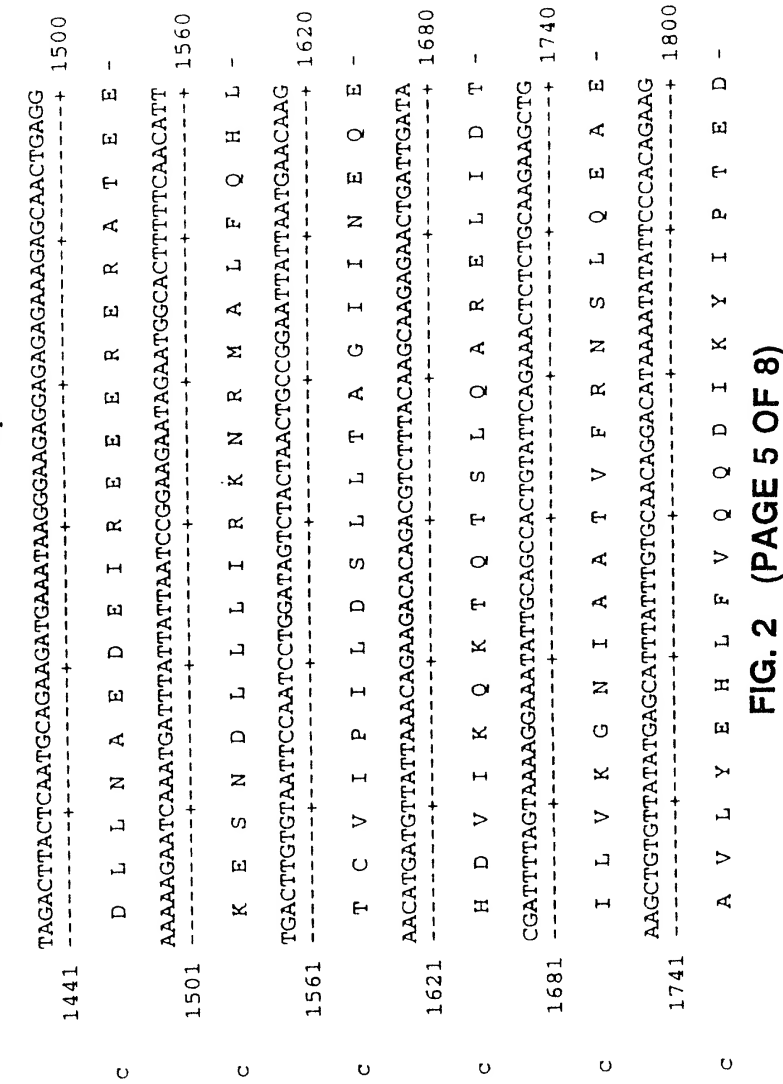


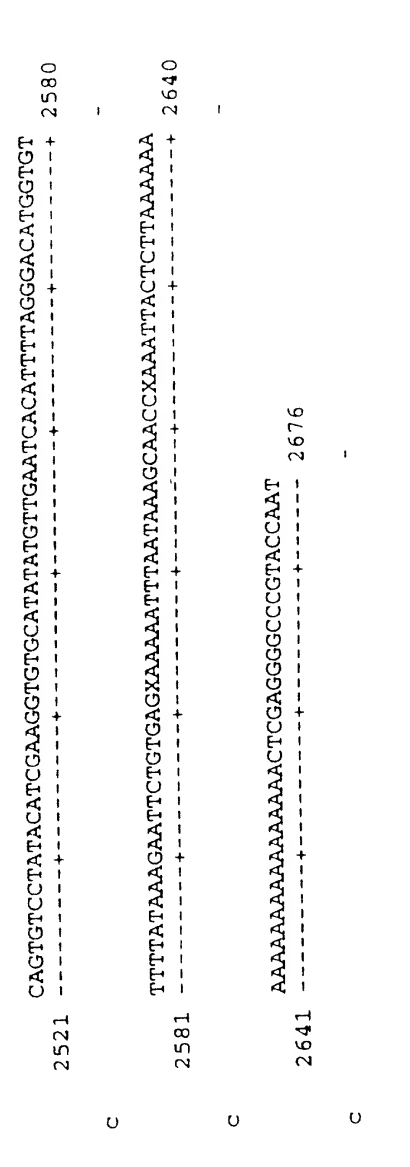
FIG. 2



	2161	TGAACATATTTTTTAGAAACTAAGAGAATGATAGGCTTTTTGTTCTTATGAAGAAAA	2220
()			ı
	2221	GAGGTAGCACTACAAACACAATATTCAATCCAAATTTCAGCATTATTGAAATTGTAAGTG	2280
U			1
	2281	AAGTAAAACTTAAGATATTTGAGTTAACCTTTTAAGAATTTTTAAATATTTTGGCATTGTAC	2340
U			ı
	2341	TAATACCGGGAACATGAAGCCAGGTGTGGTGGTATGTACCTGTAGTCCCAGGCTGAGGCA	2400
U			ŧ
	2401	AGAGAATTACTTGAGCCCAGGAGTTTGAATCCATCCTGGGCAGCATACTGAGACCCTGCC	2460
U			1
	, , , , , , , , , , , , , , , , , , ,	TTTAAAAACXAACAGXACCAAAXCCAAACACCAGGGACACATTTCTCTGTCTTTTTGAT	2520
C	1077	FIG. 2 (PAGE 7 OF 8)	ı

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## **HUMAN hiap-1**



(PAGE 8 OF 8)

FIG. 2

SEQ ID NO:7	TTAGGTTACCTGAAAGAGTTACTACAACCCCAAAGAGTTGTGTTCTAAGTAGTATCTTGG 1+++++++-
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	TAATTCAGAGAGATACTCCTACCTGAATATAAACTGAGATAAATCCAGTAAAGAAAG
ઌ	
	TGTAGTAAATTCTACATAAGAGTCTATCATTGATTTCTTTTTGTGGTGGAAATCTTAGTT 121++++++
rd	•
	CATGTGAAGAAATTTCATGTGAATGTTTTAGCTATCAAACAGTACTGTCACCTACTCATG 181++++++-240
ro	Ι ΣΣ
	CACAAAACTGCCTCCCAAAGACTTTTCCCAGGTCCCTCGTATCAAACATTAAGAGTATA 241++++++ 300
SEQ ID NO:8 a	HKTASQRLFPGPSYQNIKSI -
	ATGGAAGATAGCACGATCTTGTCAGATTGGACAAACAACAACAAAAAAAA
ro	MEDSTILSDWTNSNKQKMKY-
	FIG. 3 (PAGE 1 OF 7)

FIG. 3 (PAGE 2 OF 7)

		AA	$\mathbf{r}\mathbf{r}\mathbf{c}$	TAG	AGC.	AGT	TGA	AGAC	CATC	TC	rrc2	TCC	BAGG	ACT	AAC	CCC	TAC	PAG	PTAC	rGC	AATTCTAGAGCAGTTGAAGACATCTCTTCATCGAGGACTAACCCCTACAGTTATGCAATG	
	721	1	1	1	+	1	!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!	+	1	; ; ;	+	! !	! ! !	1	+		; 	+	ì	 	+ !	780
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	781		TAC	TGA	AGA -+-	AAGAAGC	CAG	CAGATTTCTT	PCT	rac(	CTAC	CCAC	rATC	TGC	3CC2	ATTA	AAC'	rttr +-	rtt(	GTC.	AGTACTGAAGAAGCCAGATTTCTTACCTACCATATGTGGCCATTAACTTTTTTTGTCACCA	840
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	841		AGA 	ATT	-+- 295	AAG	TCAGAATTGGCAAGAGCTGGTTTT	TGG7	rtr	rta'	rta:	TATAT.	AGGACCTGGAGA	ACC7	1991 +-	AGAC	TAG	GGTAGC +	AGC(	CTGC	TCAGAATTGGCAAGAGCTGGTTTTTTTTATATAGGACCTGGAGATAGGGTAGCCTTGCTTT	006
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	961		AGGCA	TTT1	TCC	CAA	AGGCATTTTCCCAACTGTCCATTT	FCC.	ATT	rrr 	GGA	AAAA	TTC	PCT2	AGA.	AAC'	rcr 	3AGG +-	GTT. 	rag	TTGGAAAATTCTCTAGAAACTCTGAGGTTTAGCATT	1020
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	1021		AAZ	TCAAATCT	GAG	CAT	GAGCATGCAGACACAT	GAC.	ACA	TGC 	AGC	CTCG -+	AAT(	GAG	AACATT -+	ATT' 	TATG	GTAC+-	CTG	)   	TCAAATCTGAGCATGCAGACACATGCAGCTCGAATGAGAACATTTATGTACTGGCCATCT	1080
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FIG. 3 (PAGE 3 OF 7)

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AACATCTTCAAAACTGTCTAAAAGAAATTGACTCTACATTGTATAAGAACTTATTGTG	A NIFKNCLKEIDSTLYKNLFV	GATAAGAATATGAAGTATATTCCAACAGAAGATGTTTCAGGTCTGTCACTGGAAGAACAA	A DKNMKYIPTEDVSGLSLEEQ	TTGAGGAGGTTGCAAGAACGAACTTGTAAAGTGTGTATGGACAAAGAAGTTTCTGTT 1921+++++++	A LRRLQEERTCKVCMDKEVSV	GTATTTATTCCTTGTGTCATCTGGTAGTATGCCAGGAATGTGCCCCCTTCTCTAAGAAAA 1981+++	A VFIPCGHLVVCQECAPSLRK	TGCCCTATTTGCAGGGTATAATCAAGGGTACTGTTCGTACATTTCTCTCTTAAAAAAAA	A CPICRGIIKGTVRTFLS*	ATAGTCTATATTTTAACCTGCATAAAAGGTCTTTAAAATATTGTTGAACACTTGAAGCCC 2101+++++++

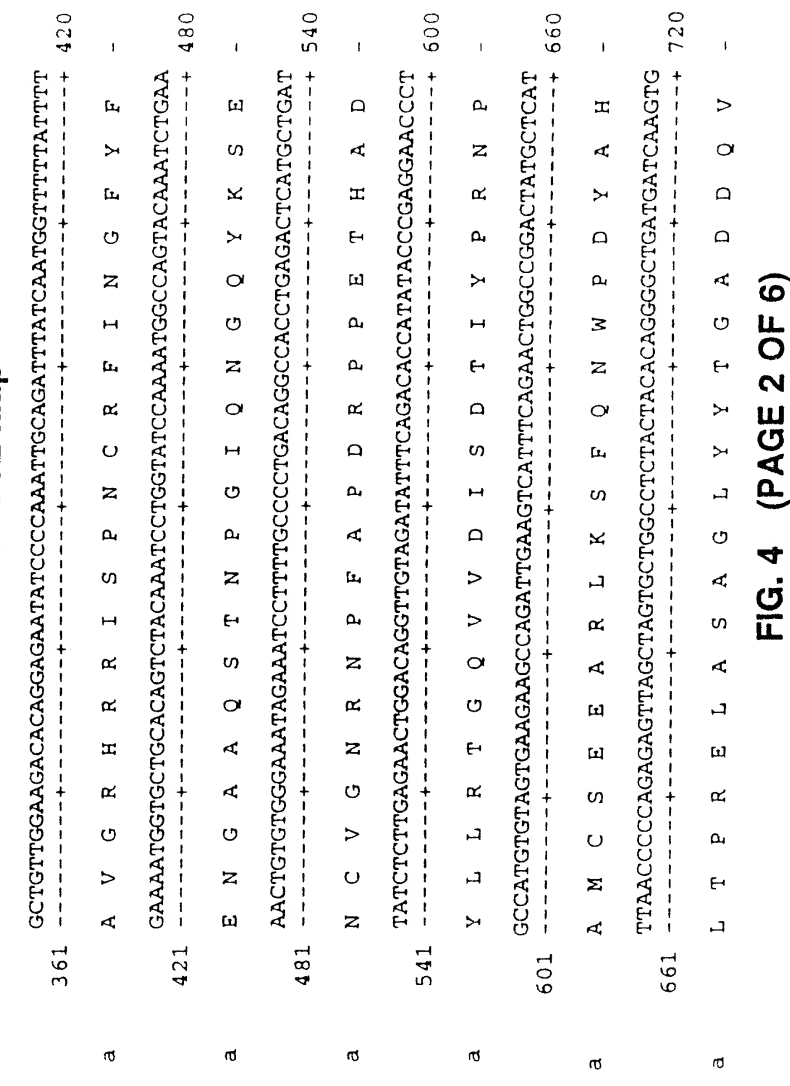
FIG. 3 (PAGE 6 OF 7)

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	2161	ATCTAAAGTAAAAGGGAATTATGAGTTTTTCAATTAGTAACATTCATGTTCTAGTCTGC 	2220
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	2221	TTTGGTACTAATAATCTTGTTTCTGAAAAGATGGTATCATATATTTTAATCTTAATCTGTT	2280
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	2281	TATTTACAAGGGAAGATTTATGTTTGGTGAACTATATTAGTATGTAT	2340
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	2341	AGTAGCGTCXCTGCTTGTTATGCATCATTTCAGGAGTTACTGGATTTGTTGTTCTTTCAG	2400
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	2401	AAAGCTTTGAAXACTAAATTATAGTGTAGAAAAGAACTGGAAACCAGGAACTCTGGAGTT	2460
て			ı
	2461	CATCAGAGTTATGGTGCCGAATTGTCTTTGGTGCTTTTCACTTGTGTTTTTAAATAAGGA	2520
ø			ſ
	2521	TTTTTCTCTTATTTCTCCCCCTAGTTTGTGAGAACATCTCAATAAAGTGCTTTAAAAAG	2580
rd		FIG. 3 (PAGE 7 OF 7)	1

	9	ı	120	1	180	1	240	ı	300	ı	360	I	
קפול חטטון	GACACTCTGCTGGGCGGCGGCCGCCCTCCTCCGGGACCTCCCCTCGGGAACCGTCGCCC		GCGGCGCTTAGTTAGGACTGGAGTGCTTGGCGCGCGAAAAGGTGGACAAGTCCTATTTTCCA 61++++++		GAGAAGATGACTTTTAACAGTTTTTGAAGGAACTAGAACTTTTGTACTTGCAGACACCAAT	M T F N S F E G T R T F V L A D T N	AAGGATGAAGAATTTGTAGAAGAGTTTTAATAGATTAAAAACATTTGCTAACTTCCCAAGT 181+++++	K D E E F N R L K T F A N F P S	AGTAGTCCTGTTTCAGCATCAACATTGGCGCGAGCTGGGTTTCTTTATACCGGTGAAGGA	SSPVSASTLARAGFLYTGEG	GACACCGTGCAATGTTTCAGTTGTCATGCGGCAATAGATAG	DTVQCFSCHAAIDRWQYGDS FIG. 4 (PAGE 1 OF E)	5
	SEQ ID NO:9	rđ		ଷ		SEQ ID NO:10 a		ิซ		rđ		rð	



721		AAT(	GCTTTTGT	TTT +	GTJ	TGTC	TGGG	GGGA +-	AAAC	CTG	AAA	AAT	TGC	GA	ACCC	TG	TGA'	rcgr +-	TGC	CTC	CAATGCTTTTGTTGTGGGGAAAACTGAAAATTGGGAACCCTGTGATCGTGCTTGGTCATCTTTGGTCAA	780
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781		AAC.	ACA	GGA +	GAC	CAC	AGGAGACACTTTCC	CCC -+-	AAT	TGC	TTT:	TT	rGTT	rrr(	3GG( - + -	) () ()	GAA	CGTT +-	TAT.	TG	GAACACAGAGACACTTTCCCAATTGCTTTTTTGTTTTGGGCCGGAACGTTAATGTTCGA	840
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841		GTG	GAATCTGGTGTGAGTTCTGAT	CTC	3GT(	GTG	AGT	TCT-+-	GAT	AGC	SAAC	. – – +	AATTTCCCAAATTCAACAAACTCTCC	4AA'	ITC.	AAC	AAA	CTO	TCC	AA(	AGTGAATCTGGTGTGAGTTCTGATAGGAATTTCCCAAATTCAACAACTCTCTCCAAGAAAT ++++++	006
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(PAGE 3 OF 6)

FIG. 4

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11.40	AAATACCTATTGGATGAGAAGGGGCAAGAATATATA +++++	TAT	GAA	CAA -+-	999	AAG	ATGAG	GAT	ATTC	FACCT/	ATA(	CAA	CATGCTAAGTGCTACCCAGGGTGC	AGG	200	CTA	GTGC -+	TAA	IGC	CA	1081	

C P M C Y T V I T F N	AAATGTCCCATGTGCTACACCGTCATTA	IVFFPCGHLATCK	ATCGTTTTTTTCCTTGTGGACATCTGGCCACTTGTAAACAGTGTGCAGAAGCAGTTGAC	
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	1921	TCATGCCTTTTGCATAAGCTTAACAAATGGAGTGTTCTGTATAAGCATGGAGATGTGATG 	1980
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	1981	GAATCTGCCCAATGACTTTAATTGGCTTATTGTAAACACGGAAAGAACTGCCCCACGCTG	2040
rd			ł
	2041	CTGGGAGGATAAAGATTGTTTTAGATGCTCACTTCTGTGTTTTTAGGATTCTGCCCATTTA	2100

FIG. 4 (PAGE 6 OF 6)

#### M-hiap-1

SEQ ID

SEQ ID

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300	TTTTCCTGTGAGCTGTACCGATTGTCCACGTATTCAGCTTTTCCCAGGGGAGTTCCTGTG	
4 7 0 4	181++++++	
0.40	GACAGCGCCTTTCTAGCCAAGCTG	
) ) 	O N W N W	F
) (	ATCCCCAGAGAAAGACTIGTCCCTTCCCCTGTCATCTCACCATGAACATGAA	
1 20	GAAGTGGGCTGCGGTATCAGCCTAGCAGTAAAACCGACCAGAAGCCATGCACAAAACTAC	
0	GAATTCCGGGAGACCTACACCCCCGGAGATCAGAGGTCATTGCTGGCGTTCAGAGCCTAG	~~

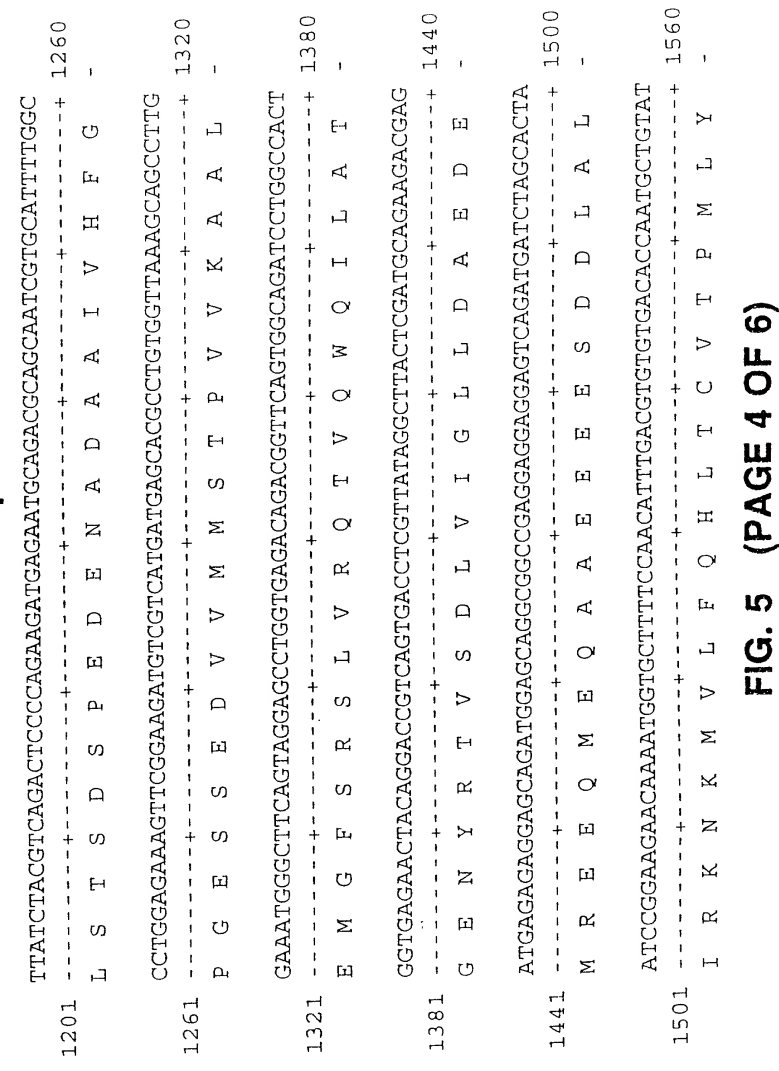
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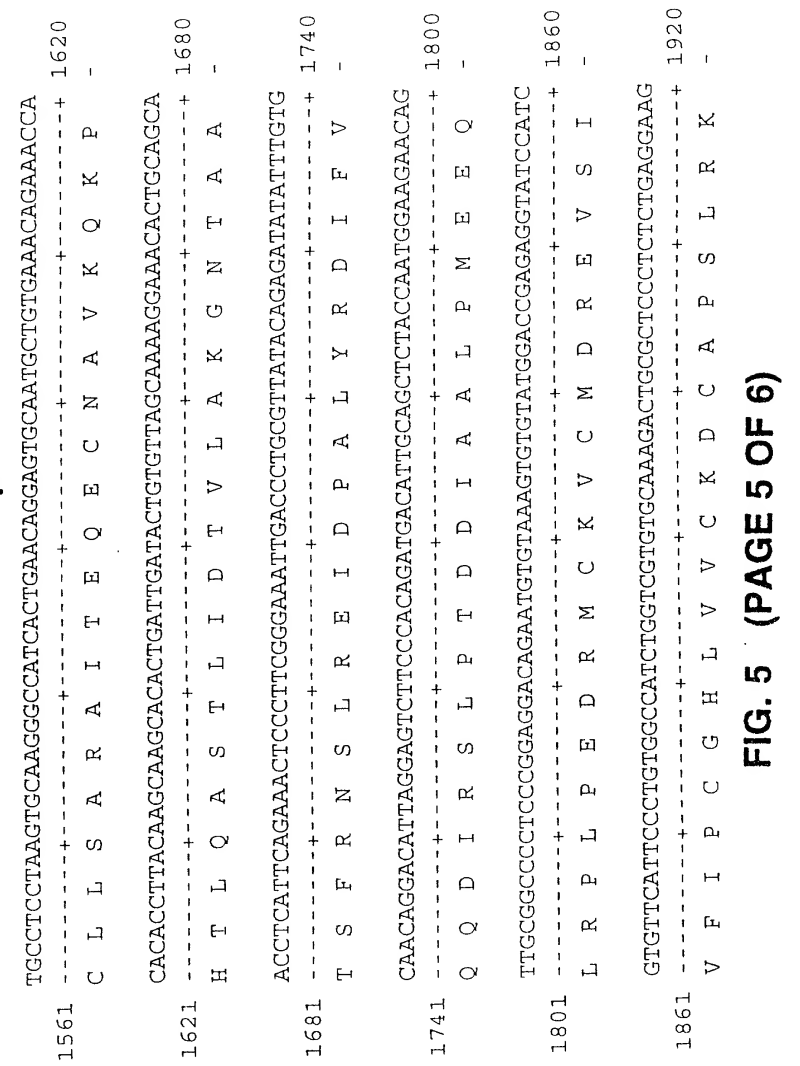
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				E G	N	Jadan Co	A	四	N	(PAGE 2 OF	6					

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707		TGC	GTG	CGA	TGG	GAZ	ACT	GAG	CAA	CTG	GGA	ACG	TAA(	GGA	TGA'	IGC	rat(	3TC	AGA	Ü	(
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							Belley	FIG.	5		(P)	(PAGE	Ш	300	OF (	9					



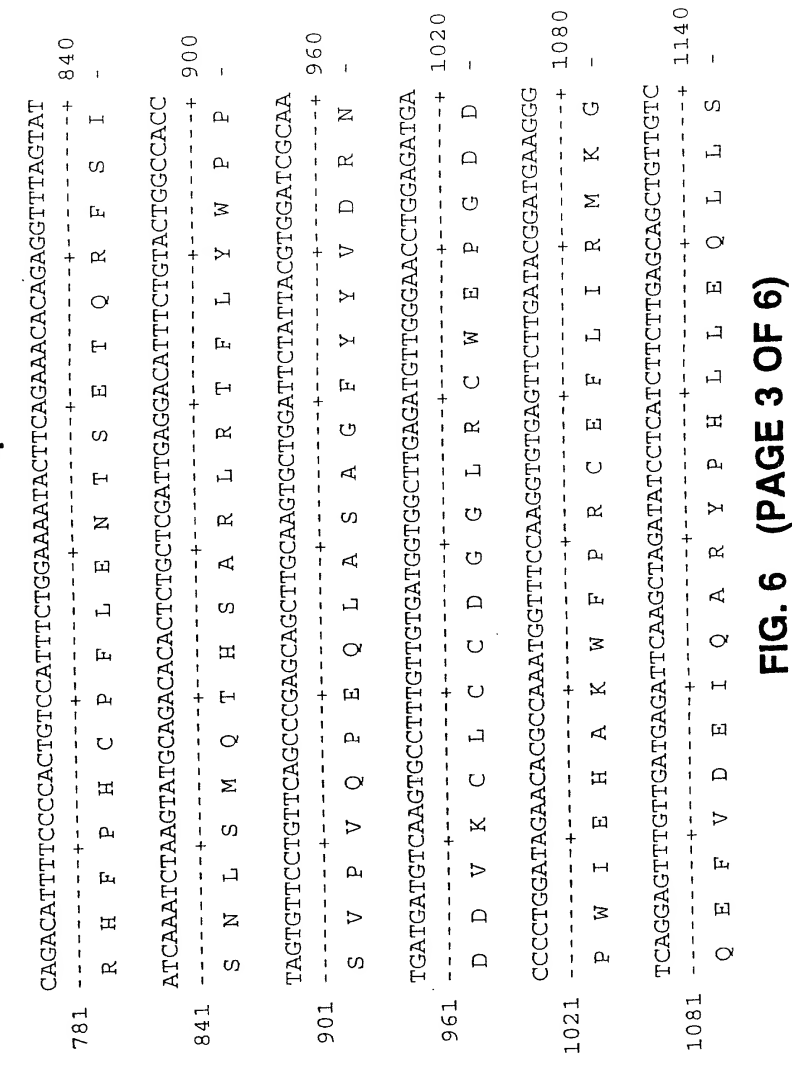


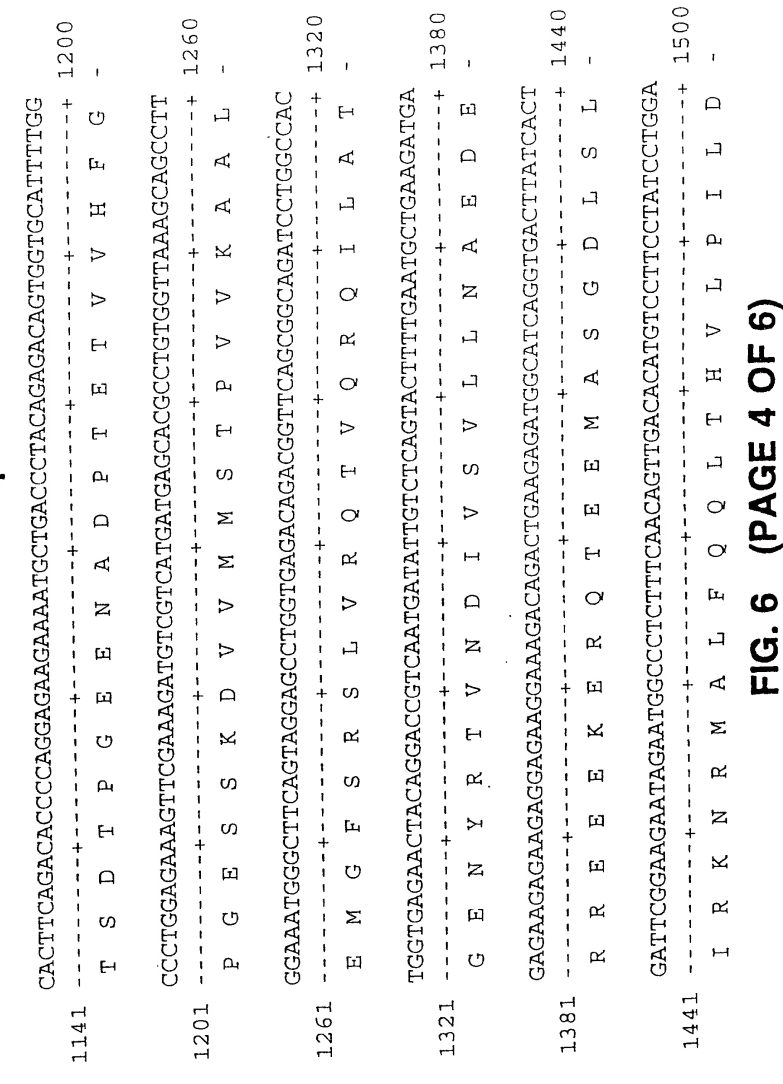
1921	TGTCCCATCTGTAGAGGGACCATCAAGGGCACAGTGCGCACATTTCTCTCTC	1980
l	CPICRGTIKGTVRTFLS*	
	CTAATGGTCCATGGCTGCAACTTCAGCCAGGAAGGTTCACTGTCACTCCCAGTTCCAT	
1981	+++	2040
2041	+1:11:+++++++++++++++++++++++++++++++++	2100
	GAAAAACTTTTGTCTGAAGTCAAGAATGAATGAATTACTTATATAATAATTTAATTGGT	
2101		2160
2161	+++++	222(
	TACTACCTGCATCTAAAGTATTCATATTCATATTTCAGATGTCATGAGAGAGGGTTT	
2221	, +	228(
	TGTTCTTGTTCCTGAAAAGCTGGTTTATCATCTGATCAGCATATACTGCGCAACGGGCAG	
2281	+	234(
	GGCTAGAATCCATGAACCAAGCTGCAAAGATCTCACGCTAAATAAGGCGGAAAGATTTGG	
2341	+1 = 1 = = =	240(
	AGAAACGAAAGGAAATTCTTTCCTGTCCAATGTATACTCTTCAGACTAATGACCTCTTCC	
2401	<u> </u>	246
2461	TATCAAGCCTTCTA FIG. 5 (PAGE 6 OF 6)	

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60	180	240	300	360	420
SEQ ID NO:41 CTGTGGTGGAGTCTATTGTCCAAGTGGTGAAACTTCATCTGGAAGTTTAAGCGGTCA  1+++++++	CACCCAAAACTTAAATGGAGAAGAGCACAATCTTGTCAAATTGGACAAAGGA	GAGCGAAGAAAAAGTTTGACTTTTCGTGTGAACTCTACCGAATGTCTACATATTC	AGCTTTTCCCAGGGGAGTTCCTGTCTCAGAGGAGTCTGGCTCGTGCTTTTATTA++++++	TACAGGTGTGAAAAGTCAAGTGCTTCTGCTGTGGCCTGATGTTGGATAACTGGAA++++++	ACAAGGGACAGTCCTGTTGAAAAGCACAGATCTATCCCAGCTGCAGCTTTGTACA+++++++ Q G D S P V E K H R Q F Y P S C S F V Q PAGE 1 OF 6)
SEQ ID NO:41 1	121 SEQ ID NO:42	181	241	301	361

780	TGCCTGTGGGAAACTGAGCAACTGGGAACCAAAGGATTATGCTATGTCAGAGCACCG +++++++  A C G K L S N W E P K D Y A M S E H R  FIG. 6 (PAGE 2 OF 6)	721
720	AGCAGAGCTGGCCAGAGCTGGCTTCTATTACATAGGGCCTGGAGACAGGGTGGCCTGTTT+++++++ A E L A R A G F Y Y I G P G D R V A C F	661
0	STEEARFL	-  
C V	GAGTACAGAAGAGGCCAGATTTCTTACTTACAGTATGTGGCCTTTAAGTTTTCTGTCACC	7
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                                                                      AGCCAACATCTTCAAAAACTCTCTGAAGGGAATTGACTCCACGTTATATGAAAACTTATT
                                   ACAGATACCCTTACAAGCAAGAGCTTATTGACACCGTTTTAGTCAAGGGAAATGCTGC
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1	CTICTIGGGATTIGGGAATTIGGGGAAAGCTTIGGAATCCAGIGATGIGGAGCTCAGAAA	( ( (
2161	++++++	0777
2221		2280
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2341		2400
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# Alignment of BIR (Baculoviral IAP Repeats) Domains

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Cp\_iap Cydia pomonella Op\_iap Orgyia pseudotsugata

Human

IAP on X chromosome two different human IAP genes

FIG. 7

xiap hiap1, hiap2 mouse homologue of human xiap gene

m-xiap Insect

diap

Drosophilia IAP gene, not clearly a homologue of xiap or hiap

note on consensus: The co

The consensus line represents amino acids or very similar amino acids which are present in 14 of the 19 BIR sequences at each position.

Capitalized residues are those that are in the consensus sequence.

68 WPwof lens rWassGFYY1 GrobeVrCaf CkweitnWwr ondheidHkr wapnchFV	. 1spe tMAknGFYY1 GrsDeVrCaf CkveimrWke gEdpaadHkk	WPnpn.ilpg alakaGFYY1 nrlDhVkCvw CngviakWek nDnafeeHkr ffPqCprV	Presspysas tharagrift degotvecre chasidrwcy gosavgrhrr ispucrfi	FPsgspvsas tlaraGFLYt GegDtVrCFs ChaavdrWqy gDsavgrHrk vsPnCrFI	Pagypyser slaraGFYYt GynDkVkCFc CglmldnWkr gDsptekEkk lyPsCrFV	FPagvpvser staraGFYYt GvnDkVkCFc CglmldnWkl gDspiqkHkq lyPsCsFI	WPdyahltpr eLAsAGLYYt GadDqVqCFc CggklknWep cDrawseHrr hfPnCfFV	WPdyahltpr elasaGlyyt GigDqVqCFc CggklknWep cDrawseRrr hfPnCfFV	WP.liflspt dlaraGFYYi GpgDrVaCFa CggklsnWep kDnamseHlr hfPkCpFI	elaragrii gpgDrVaCFa CggklsnWep kDdamseHrr	qLArAGFYal GegDkVkCFh CgggltdWkp sEdpwdqHak	qlaragryal GegDkvkCrb	WPssvlvnpe qLAsAGFYYv GnsDdVkCFc CdgglrcWes gDdpwvqHak wfPrCeYl	CdgglrcWes gDdpwveHak	kqrpe elaeaGFFYt GqgDktrCFc	ukqrpe qMAdAGFFYt GygDntkCFy	WPisniqpas alaqaGlYYq kigDqVrCFh CniglrsWqk eDepwieHak wsPkCqFV	Inapvsae divangif GtwmeaeCdf	WPWLA-AGFYY- GD-V-CF- CWDHP-C-FV
1 kaarronytn W		eanRLvTFkd W	efnRLkTFan F	efnRikIran F	elyRMsTYst F	elyRMsTYst F	eeaRLksfqn W		enaRtlTFqt W	eeaRFlTYhm W	yearivfff W	yearifrrgt w	baaRFkTFfn W	baaRMrTFmy W	eaarlrTFae W	eaaRvksFhn W	vdaRirTFtd W	esvRlaTFge P	RL-TF V
0 4 1 1	Cp_iap-1	diap-2	m-xiap-1	xiap-1	hiap1-1	hiap2-1	m-xiap-2	xiap-2	hiap1-2	hiap2-2	m-xtap-3	xtap-3	hiapi-3	hiap2-3	op_iap-2	Cp_iap-2	diap-3	diap-1	Consensus
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SEQ ID NO:12	cp-iap	• • • • • • • •	• • • • • • • • • • •			
SEQ ID NO:13	diap		• • • • • • • • • •	mtelgMelEs	VRLaTFgeWP	lnaPVSaedL
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SEQ ID NO:4	xiap	mtinsie	gsktcvpadi	nkeeEFveEF	nRLkTFanFP	sgsPVSastL
SEQ ID NO:6	hiap1	mnivensifl	snlmksantf	elkyDLscEL	yRMsTYstFP	aqv?VSersL
SEQ ID NO:8	. hiap2			kmkyDFscEL		
SEQ ID NO:44	consensus			*	-RL-TFFP	_
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·	cp-lap				• • • • • • • • •	
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·				aavDrWqyGD	_	,
			•	lmlDnWkrGD	-	,
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		•	•	lmlDnWklGD	• • •	1 •
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Table of the state	consensus	2	D-SD	M	EEARL-TF	WPL-
terror state acting light		•	BIR 2	•		
		201			<del></del>	250
	cp-lap	PetMAknGFY	YlGrsDeVrC	afCkveimrW	kegEdpaaDH	kkwaPqCPFV
	dlap	PoalakaGFY	YlnrlDhVkC	vwCnGviakW	EknDnAfeEH	kRfFPqCPrV
•	m-xiap	Prelasagly	YtGadDqVqC	FcCGGKLkNW	EPcDrAwsEH	rRHFPnCfFV
	xiap	PrelasaGLY	YtGigDqVqC	FCCGGKLKNW	EPCDrAwSEH	rRHFPnCfFV
	hiap1	PtDLArAGFY	YiGpgDrVaC	Facggklsnw	EPkDnAmSEH	lRHFPkCPFI
	hiap2	PEELARAGEY	YiGpgDrVaC	Facggklanw	EPkDdAmSEH	rRHF?nCPF1
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	cp-ian	. kaidvaasiv	ttnnighttt	hdtiigPahP	kyAheaARuk	sFhnWPrcmk
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	hiap1	. TATHTHTTO	engladear)	tvsNl	. smothaarf	TFfnWPsSvl
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Fig. 8 (page 1 of 3)

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	orpEQMAdAG	FFYtGyGDnt	KCFVCdGGLk	dWepeDvPWe	CHVIMEGICA
•	cpasalhqAG	LYYqkiGDqV		swakeDEPWf	eHAKWs PkCq
	VnkECLArAG	•	KCFhCgGGLt	cWkpsEDPWd	CHAKCYPgCk
_ ·	VakEQLAFAG	FYalGeGDkV		dWkpsEDPWe	QHAKWYPqCk
•	VabEQLASAG	FYYvGnsDdV		CHESCODPWV	CHAKWEPTCe
	VapEQLASAG	FYYWGEnDdV		cwesqDDPWv	eHAKWEFICe
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cp-lap	acVLpge.				
diap	. adVLmcea	pakeAltIGi	dggvVrnaiq	rKllssGcaF	
m-xiap	kiddtifqnP	mvqeAirMGF	sikdlKktme	eKIqtsGssY	lslevilaDL
xiap	riDdtifqnP	mVqeAirMGF	sikdlKkime	eKIqisGsnY	kslevlVaDL
hiap1	seDaIMmntP	vInaAveMGF	spayAxdoad		
hiap2	seDaVMmntP	vVksAleMGF	urclyKdraj	sKIlttGenY	ktvndiVsal
consensus	5-V9	-VAMGF	VK	-KIGY	1V-DL
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cp-lap					
dlap	ficagagaal	Evreppe			
m-xiap	vsAqkDnieD				
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diap	sniskitdei	dkwanacbud	nlSlowenRg	LkDarLCKVC	LDeEVgVVFl
m-xlap	. ,		aists QLRR	LqEEkLsKIC	MDrnIaIVFf
xlap			ersteects	LqEEkLCKIC	MDrnlalVFV
hiap1	lyehlfraga	ikyiptedvs	dlpvEEOLRR	LpEErtCKVC	MDkEVsIVFI
hlap2	lyknliwdkn	mkylptedvs	glslemeQLRR	LqEExtCKVC	MDkEVsVVFI
consensus			S-EEQLRR	I-EE-ICK-C	MD-EVVF-
	601			635	
cp-iap	PCGHvVaCak	CALSVCKCPM	CRKIVtsvlk	VYFS.	
diap	PCGHLatCnq	CApSVanCPM	CRadIkgivi	tfls*	
m-xiap	PCGHLatCkq	CAeaVdKCPM	CytVItingk	1FMS™	
xiap	_	CAeaVdKCPM	į .		
hlap1		CApSlrKCPi	1		
hiap2	_	CApSlrKCPi			
consensus	PCGHLV-C	CA-SV-KCPM		-FLS-	

## Alignment of RZF (Ring Zinc Finger) Domains

Baculovirus

Cydia pomonella Cp\_iap Op\_ap

Orgyia pseudotsugata

Human

xiap

IAP on X chromosome

two different human IAP genes hiap1, hiap2

Mouse

mouse homologue of human xiap gene

Insect

diap

m-xiap

Drosophilia IAP gene, not clearly a homologue of xiap or hiap

note on consensus:

The consensus line represents amino acids or very similar amino acids which are present in 6 of the 7 RZF sequences at each position.

Capitalized residues are those that are in the consensus sequence.

tckvcMdkev tCKVCMdkev lsKICMdrni EqlrriqEer Eqitripser Eqitriquek Eqitriquek n-xiad hiap1 hiap2 xiap NO:33 NO:32 NO:34 NO:35 H A SEQ SEQ SEQ

Cp\_iap Op\_iap diap SEQ ID NO: 1 ID NO:36 ID NO:37 SEQ SEQ

Consensus

ttcpvc dkCPmC ancemc vacakcalsv vtckqcAeav atCnqCApev vacgkchagv --C--CA-**tvcrvpcghy** LVCFVPCGHV alvrvPCGH1 -V-F-PCGHgvyflpcgh1 1CKICLGack LCKVCLdeev 1CKICyveec -CKICM---aveaevaDdr EenrglkDar Ekepqvedsk

rkcric dkcpmc dkcpmc

atckqcAeav

w CkdCApsl

BIVFIPCGET

AIVF FPCGH1

rkcPic

svvripcGH1 vvcqeCApel

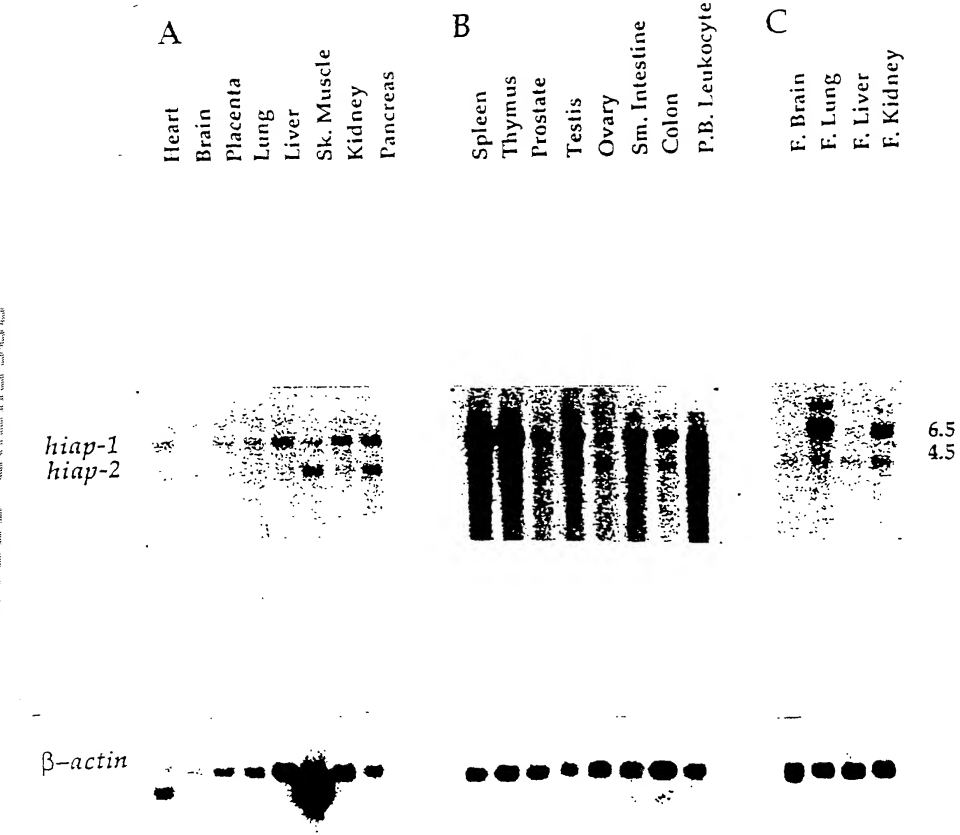
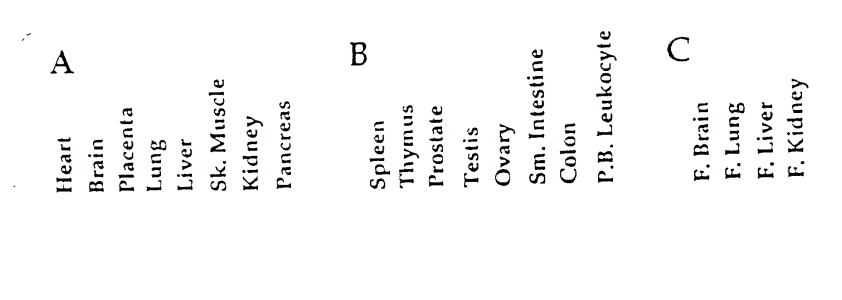


FIG. 10







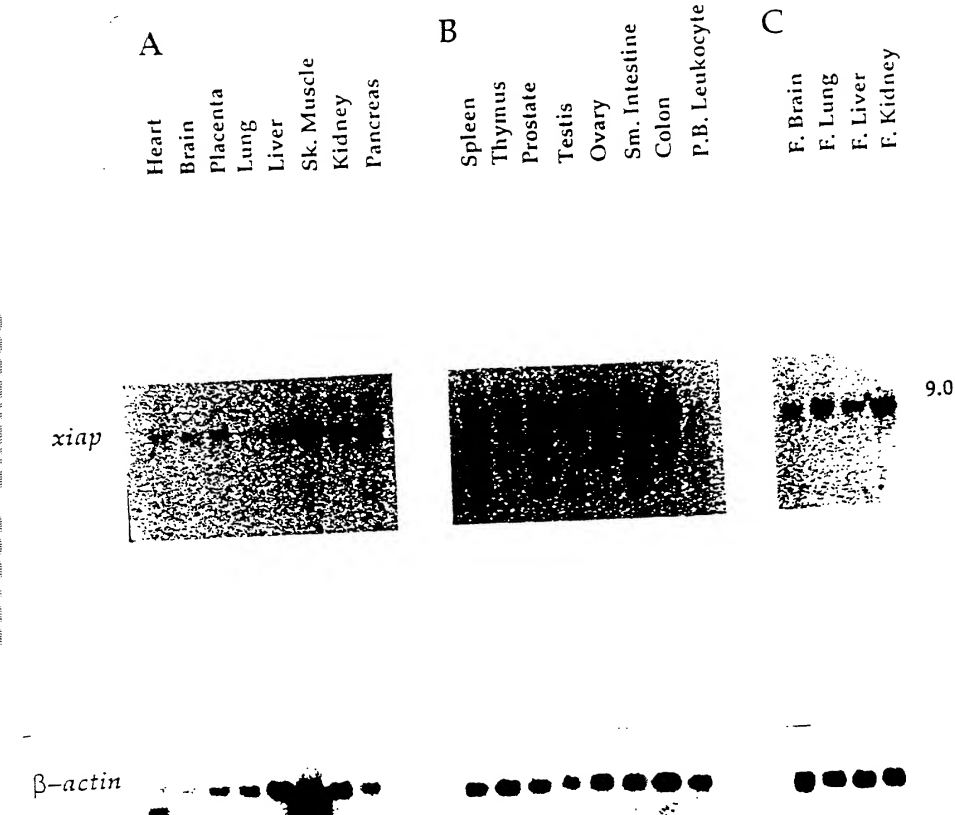
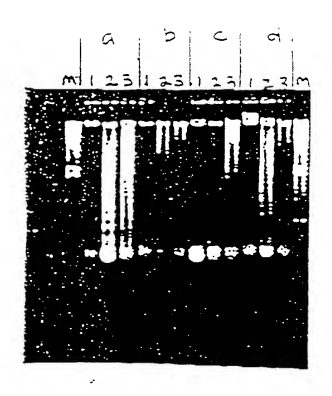


FIG. 12



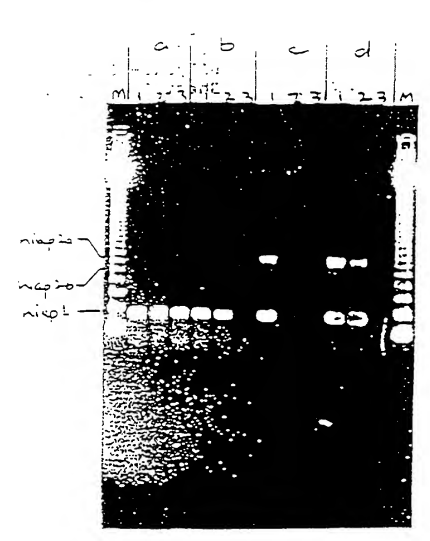


Fig. 13A and 13B

Fig. 14A - D

xiap

#### COMBINED DECLARATION AND POWER OF ATTORNEY

As a below named inventor, I hereby declare that:

My residence, post office address and citizenship are as stated below next to my name,

first and joint inventor which a patent is sough PROBES AND DETERMINED IS attached	r (if plural names are list that on the invention entite ECTIONS METHODS, thereto.	ele inventor (if only one name is listed below) or an original, ted below) of the subject matter which is claimed and for eled MAMMALIAN IAP GENE FAMILY, PRIMERS, the specification of which application Serial No. 08/576,956 and was amended on
was filed of	December 22, 1995 as A	application Serial No. 08/370,330 and was amended on
——· □ was describ	ed and claimed in PCT I	nternational Application No.
filed on	and a	s amended under PCT Article 19 on
I hereby state specification, including	that I have reviewed and the claims, as amended	l understand the contents of the above-identified by any amendment referred to above.
	the duty to disclose all it 37, Code of Federal Reg	information I know to be material to patentability in ulations, §1.56(a).
application(s) listed be disclosed in the prior United States Code, § patentability as defined	clow and, insofar as the s United States application 112, I acknowledge the d d in Title 37, Code of Fe	35, United States Code, §120 of any United States subject matter of each of the claims of this application is not in the manner provided by the first paragraph of Title 35, uty to disclose all information I know to be material to deral Regulations, §1.56(a) which became available between national or PCT international filing date of this application:
U.S. SERIAL NO.	FILING DATE	STATUS  The diag Discussion Discu
08/511.485	August 4, 1995	■ Pending □ Issued □ Abandoned

I hereby appoint the following attorneys and/or agents to prosecute this application and to transact all business in the Patent and Trademark Office connected therewith: Paul T. Clark, Reg. No. 30,162 and Kristina Bieker-Brady, Reg. No. 39,109, William E. Booth, Reg. No. 28,933; Barry E. Bretschneider, Reg. No. 28,055; John W. Freeman, Reg. No. 29,066; Timothy A. French, Reg. No. 30,175; Alan H. Gordon, Reg. No. 26,168; John F. Land, Reg. No. 29,554; John B. Pegram, Reg. No. 25,198; Rene D. Tegtmeyer, Reg. No. 33,567; Hans R. Troesch, Reg. No. 36,950; Dorothy P. Whelan, Reg. No. 33,814; Charles C. Winchester, Reg. No. 21,040.

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Address all correspondence to <u>Kristina Bieker-Brady</u>, Fish & Richardson P.C., 225 Franklin Street, Boston, MA 02110-2804.

I hereby declare that all statements made herein of my own knowledge are true and that all statements made on information and belief are believed to be true; and further that these statements were made with the knowledge that willful false statements and the like so made are punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that such willful false statements may jeopardize the validity of the application or any patents issued thereon.

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Inventor's Signature: 5th Baid Date: Ma 18/96
Residence Address: Ottawa, Ontario, Canada
Citizen of: Canada
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167882.B11

#### SEQUENCE LISTING

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Cys Xaa Phe Val
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100

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Pro Asn Cys Phe Phe Val Leu Gly Arg Asn Leu Asn Ile Arg Ser Glu
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Gly Gly Leu Thr Asp Trp Lys Pro Ser Glu Asp Pro Trp Glu Gln His
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Glu Tyr Ile Asn Asn Ile His Leu Thr His Ser Leu Glu Glu Cys Leu
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Gly Ser Asn Tyr Lys Ser Leu Glu Val Leu Val Ala Asp Leu Val Asn
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Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp Lys Arg Gly Asp
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Phe Pro Ser Ser Val Thr His Ser Thr His Ser Leu Leu Pro Gly Thr
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Glu Asn Ser Gly Tyr Phe Arg Gly Ser Tyr Ser Asn Ser Pro Ser Asn
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Pro Val Asn Ser Arg Ala Asn Gln Glu Phe Ser Ala Leu Met Arg Ser
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Gln Thr Trp Pro Leu Thr Phe Leu Ser Pro Thr Asp Leu Ala Arg Ala
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Gly Phe Tyr Tyr Ile Gly Pro Gly Asp Arg Val Ala Cys Phe Ala Cys
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Tyr Leu Ile Arg Ile Lys Gly Gln Glu Phe Ile Arg Gln Val Gln Ala
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Arg Ala Gly Phe Leu Tyr Thr Gly Glu Gly Asp Thr Val Gln Cys Phe
Ser Cys His Ala Ala Ile Asp Arg Trp Gln Tyr Gly Asp Ser Ala Val
Gly Arg His Arg Arg Ile Ser Pro Asn Cys Arg Phe Ile Asn Gly Phe
Tyr Phe Glu Asn Gly Ala Ala Gln Ser Thr Asn Pro Gly Ile Gln Asn
                                105
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Gly Gln Tyr Lys Ser Glu Asn Cys Val Gly Asn Arg Asn Pro Phe Ala
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                            120
Pro Asp Arg Pro Pro Glu Thr His Ala Asp Tyr Leu Leu Arg Thr Gly
                        135
                                            140
Gln Val Val Asp Ile Ser Asp Thr Ile Tyr Pro Arg Asn Pro Ala Met
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                    150
Cys Ser Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr
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                                    170
Ala His Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Thr
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Gly Ala Asp Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys
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Asn Trp Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe
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                        215
Pro Asn Cys Phe Phe Val Leu Gly Arg Asn Val Asn Val Arg Ser Glu
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Ser Gly Val Ser Ser Asp Arg Asn Phe Pro Asn Ser Thr Asn Ser Pro
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Arg Asn Pro Ala Met Ala Glu Tyr Glu Ala Arg Ile Val Thr Phe Gly
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Thr Trp Ile Tyr Ser Val Asn Lys Glu Gln Leu Ala Arg Ala Gly Phe
                            280
Tyr Ala Leu Gly Glu Gly Asp Lys Val Lys Cys Phe His Cys Gly Gly
                        295
Gly Leu Thr Asp Trp Lys Pro Ser Glu Asp Pro Trp Asp Gln His Ala
                    310
                                        315
Lys Cys Tyr Pro Gly Cys Lys Tyr Leu Leu Asp Glu Lys Gly Gln Glu
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Tyr Ile Asn Asn Ile His Leu Thr His Pro Leu Glu Glu Ser Leu Gly
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Arg Thr Ala Glu Lys Thr Pro Pro Leu Thr Lys Lys Ile Asp Asp Thr
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Ile Phe Gln Asn Pro Met Val Gln Glu Ala Ile Arg Met Gly Phe Ser
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Phe Lys Asp Leu Lys Lys Thr Met Glu Glu Lys Ile Gln Thr Ser Gly
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                    390
Ser Ser Tyr Leu Ser Leu Glu Val Leu Ile Ala Asp Leu Val Ser Ala
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Gln Lys Asp Asn Thr Glu Asp Glu Ser Ser Gln Thr Ser Leu Gln Lys
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Asp Ile Ser Thr Glu Glu Gln Leu Arg Arg Leu Gln Glu Glu Lys Leu
                                                445
Ser Lys Ile Cys Met Asp Arg Asn Ile Ala Ile Val Phe Pro Cys
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Gly His Leu Ala Thr Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys
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<210> 11
<211> 67
<212> PRT
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<213> Orgyia pseudotsugata

<400> 11

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<400> 12

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Lys Ser Phe His Asn Trp Pro Arg Cys Met Lys Gln Arg Pro Glu Gln
                            120
Met Ala Asp Ala Gly Phe Phe Tyr Thr Gly Tyr Gly Asp Asn Thr Lys
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Cys Phe Tyr Cys Asp Gly Gly Leu Lys Asp Trp Glu Pro Glu Asp Val
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Pro Trp Glu Gln His Val Arg Trp Phe Asp Arg Cys Ala Tyr Val Gln
Leu Val Lys Gly Arg Asp Tyr Val Gln Lys Val Ile Thr Glu Ala Cys
                                185
Val Leu Pro Gly Glu Asn Thr Thr Val Ser Thr Ala Ala Pro Val Ser
                            200
Glu Pro Ile Pro Glu Thr Lys Ile Glu Lys Glu Pro Gln Val Glu Asp
                        215
    210
Ser Lys Leu Cys Lys Ile Cys Tyr Val Glu Glu Cys Ile Val Cys Phe
                                        235
                    230
Val Pro Cys Gly His Val Val Ala Cys Ala Lys Cys Ala Leu Ser Val
                                    250
Asp Lys Cys Pro Met Cys Arg Lys Ile Val Thr Ser Val Leu Lys Val
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Tyr Phe Ser
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<213> Drosophila melanogaster
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Asn Gly Phe Phe Ala Thr Gly Lys Trp Leu Glu Ala Glu Cys His Phe
                            40
Cys His Val Arg Ile Asp Arg Trp Glu Tyr Gly Asp Gln Val Ala Glu
                                             60
Arg His Arg Arg Ser Ser Pro Ile Cys Ser Met Val Leu Ala Pro Asn
                                        75
His Cys Gly Asn Val Pro Arg Ser Gln Glu Ser Asp Asn Glu Gly Asn
Ser Val Val Asp Ser Pro Glu Ser Cys Ser Cys Pro Asp Leu Leu Leu
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 Ser
 Val
 Asp
 Ser
 Pro
 Glu
 Ser
 Cys
 Ser
 Cys
 Pro
 Asp
 Leu
 Leu
 Leu
 Leu

 Glu
 Ala
 Asn
 Arg
 Leu
 Val
 Thr
 Phe
 Lys
 Asp
 Trp
 Pro
 Asn
 Pro
 Asn
 Ile

 Thr
 Pro
 Gln
 Ala
 Leu
 Ala
 Lys
 Ala
 Gly
 Phe
 Tyr
 Pro
 Asn
 Pro
 Asn
 Arg
 Leu

 Asp
 His
 Val
 Lys
 Ala
 Lys
 Ala
 Gly
 Phe
 Tyr
 Tyr
 Leu
 Asn
 Arg
 Leu

 Asp
 His
 Val
 Lys
 Val
 Trp
 Cys
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 Gly
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 Ala
 Lys
 Trp
 Ile

 Lys
 Asp
 Asp
 Asp
 Ala
 Phe
 Glu
 His
 Lys
 Arg
 Phe
 Phe
 Phe
 Phe
 Phe
 Ile
 Ile
 Ala
 Thr
 Glu
 Lys
 Asp
 A

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Leu Asp Glu Leu Gly Ile Gln Pro Thr Thr Leu Pro Leu Arg Pro Lys
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Tyr Ala Cys Val Asp Ala Arg Leu Arg Thr Phe Thr Asp Trp Pro Ile
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Ser Asn Ile Gln Pro Ala Ser Ala Leu Ala Gln Ala Gly Leu Tyr Tyr
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                                        235
Gln Lys Ile Gly Asp Gln Val Arg Cys Phe His Cys Asn Ile Gly Leu
                                    250
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Arg Ser Trp Gln Lys Glu Asp Glu Pro Trp Phe Glu His Ala Lys Trp
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Ser Pro Lys Cys Gln Phe Val Leu Leu Ala Lys Gly Pro Ala Tyr Val
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Ser Glu Val Leu Ala Thr Thr Ala Ala Asn Ala Ser Ser Gln Pro Ala
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Thr Ala Pro Ala Pro Thr Leu Gln Ala Asp Val Leu Met Asp Glu Ala
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                                        315
Pro Ala Lys Glu Ala Leu Thr Leu Gly Ile Asp Gly Gly Val Val Arg
Asn Ala Ile Gln Arg Lys Leu Leu Ser Ser Gly Cys Ala Phe Ser Thr
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Leu Asp Glu Leu Leu His Asp Ile Phe Asp Asp Ala Gly Ala Gly Ala
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Ala Leu Glu Val Arg Glu Pro Pro Glu Pro Ser Ala Pro Phe Ile Glu
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Pro Cys Gln Ala Thr Thr Ser Lys Ala Ala Ser Val Pro Ile Pro Val
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Ala Asp Ser Ile Pro Ala Lys Pro Gln Ala Ala Glu Ala Val Ser Asn
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Ile Ser Lys Ile Thr Asp Glu Ile Gln Lys Met Ser Val Ser Thr Pro
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Asn Gly Asn Leu Ser Leu Glu Glu Glu Asn Arg Gln Leu Lys Asp Ala
                            440
Arg Leu Cys Lys Val Cys Leu Asp Glu Glu Val Gly Val Val Phe Leu
                        455
                                            460
Pro Cys Gly His Leu Ala Thr Cys Asn Gln Cys Ala Pro Ser Val Ala
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Asn Cys Pro Met Cys Arg Ala Asp Ile Lys Gly Phe Val Arg Thr Phe
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<210> 14

Leu Ser

<211> 67

<212> PRT

<213> Cydia pomonella

<400> 14

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Ser Pro Glu Thr Met Ala Lys Asn Gly Phe Tyr Tyr Leu Gly Arg Ser 20 25 30

Asp Glu Val Arg Cys Ala Phe Cys Lys Val Glu Ile Met Arg Trp Lys 35 40 45

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Pro Phe Val
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<213> Homo sapiens
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Thr Pro Gln Ala Leu Ala Lys Ala Gly Phe Tyr Tyr Leu Asn Arg Leu
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Asp His Val Lys Cys Val Trp Cys Asn Gly Val Ile Ala Lys Trp Glu
Lys Asn Asp Asn Ala Phe Glu Glu His Lys Arg Phe Phe Pro Gln Cys
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Pro Arg Val
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<210> 16
<211> 68
<212> PRT
<213> Mus musculus
<400> 16
Glu Phe Asn Arg Leu Lys Thr Phe Ala Asn Phe Pro Ser Ser Ser Pro
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Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu
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Gly Asp Thr Val Gln Cys Phe Ser Cys His Ala Ala Ile Asp Arg Trp
Gln Tyr Gly Asp Ser Ala Val Gly Arg His Arg Arg Ile Ser Pro Asn
Cys Arg Phe Ile
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<210> 17
<211> 68
<212> PRT
<213> Homo sapiens
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Glu Phe Asn Arg Leu Lys Thr Phe Ala Asn Phe Pro Ser Gly Ser Pro
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Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu
Gly Asp Thr Val Arg Cys Phe Ser Cys His Ala Ala Val Asp Arg Trp
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Gln Tyr Gly Asp Ser Ala Val Gly Arg His Arg Lys Val Ser Pro Asn
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Cys Arg Phe Ile
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<211> 68
<212> PRT
<213> Homo sapiens
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Glu Leu Tyr Arg Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro
Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val
Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp
Lys Arg Gly Asp Ser Pro Thr Glu Lys His Lys Lys Leu Tyr Pro Ser
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                        55
Cys Arg Phe Val
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<210> 19
<211> 68
<212> PRT
<213> Homo sapiens
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Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val
Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp
Lys Leu Gly Asp Ser Pro Ile Gln Lys His Lys Gln Leu Tyr Pro Ser
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Cys Ser Phe Ile
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<211> 68
<212> PRT
<213> Mus musculus
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Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr Ala His
Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr Gly Ala
Asp Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp
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Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn
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Cys Phe Phe Val
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<210> 21
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<212> PRT
<213> Homo sapiens
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Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr Gly Ile
Gly Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp
Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn
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Cys Phe Phe Val
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<213> Homo sapiens
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Ser Pro Thr Asp Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly
Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu
Pro Lys Asp Asn Ala Met Ser Glu His Leu Arg His Phe Pro Lys Cys
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Pro Phe Ile
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<210> 23
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<213> Homo sapiens
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Glu Glu Ala Arg Phe Leu Thr Tyr His Met Trp Pro Leu Thr Phe Leu
Ser Pro Ser Glu Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly
Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu
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Pro Lys Asp Asp Ala Met Ser Glu His Arg Arg His Phe Pro Asn Cys
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Pro Phe Leu
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<210> 24
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<212> PRT
<213> Mus musculus
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Tyr Glu Ala Arg Ile Val Thr Phe Gly Thr Trp Ile Tyr Ser Val Asn
Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp
Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro
Ser Glu Asp Pro Trp Asp Gln His Ala Lys Cys Tyr Pro Gly Cys Lys
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Tyr Leu
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<213> Homo sapiens
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Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp
Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro
Ser Glu Asp Pro Trp Glu Gln His Ala Lys Trp Tyr Pro Gly Cys Lys
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Tyr Leu
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<213> Homo sapiens
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His Ala Ala Arg Phe Lys Thr Phe Phe Asn Trp Pro Ser Ser Val Leu
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Ser Asp Asp Val Lys Cys Phe Cys Cys Asp Gly Gly Leu Arg Cys Trp
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Glu Ser Gly Asp Asp Pro Trp Val Gln His Ala Lys Trp Phe Pro Arg
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Cys Glu Tyr Leu
65
<210> 27
<211> 68
<212> PRT
<213> Homo sapiens
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His Ala Ala Arg Met Arg Thr Phe Met Tyr Trp Pro Ser Ser Val Pro
Val Gln Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Arg
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Glu Ser Gly Asp Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg
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Cys Glu Phe Leu
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<210> 28
<211> 68
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<213> Orgyia pseudotsugata
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Gln Arg Pro Glu Glu Leu Ala Glu Ala Gly Phe Phe Tyr Thr Gly Gln
Gly Asp Lys Thr Arg Cys Phe Cys Cys Asp Gly Gly Leu Lys Asp Trp
Glu Pro Asp Asp Ala Pro Trp Gln Gln His Ala Arg Trp Tyr Asp Arg
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Cys Glu Tyr Val
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<210> 29
<211> 68
<212> PRT
<213> Cydia pomonella
<400> 29
Glu Ala Ala Arg Val Lys Ser Phe His Asn Trp Pro Arg Cys Met Lys
Gln Arg Pro Glu Gln Met Ala Asp Ala Gly Phe Phe Tyr Thr Gly Tyr
Gly Asp Asn Thr Lys Cys Phe Tyr Cys Asp Gly Gly Leu Lys Asp Trp
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<213> Homo sapiens

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Glu Pro Glu Asp Val Pro Trp Glu Gln His Val Arg Trp Phe Asp Arg
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Cys Ala Tyr Val
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<210> 30
<211> 68
<212> PRT
<213> Drosophila melanogaster
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Gln Pro Ala Ser Ala Leu Ala Gln Ala Gly Leu Tyr Tyr Gln Lys Ile
Gly Asp Gln Val Arg Cys Phe His Cys Asn Ile Gly Leu Arg Ser Trp
Gln Lys Glu Asp Glu Pro Trp Phe Glu His Ala Lys Trp Ser Pro Lys
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Cys Gln Phe Val
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<213> Drosophila melanogaster
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Glu Ser Val Arg Leu Ala Thr Phe Gly Glu Trp Pro Leu Asn Ala Pro
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Val Ser Ala Glu Asp Leu Val Ala Asn Gly Phe Phe Gly Thr Trp Met
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Glu Ala Glu Cys Asp Phe Cys His Val Arg Ile Asp Arg Trp Glu Tyr
Gly Asp Leu Val Ala Glu Arg His Arg Arg Ser Ser Pro Ile Cys Ser
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Met Val
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<213> Homo sapiens
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Cys Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys
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<212> PRT
<213> Homo sapiens
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Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys Pro Met Cys
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<210> 35
<211> 46
<212> PRT
<213> Homo sapiens
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Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys Pro Met Cys
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<213> Drosophila melanogaster
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Asp Glu Glu Val Gly Val Val Phe Leu Pro Cys Gly His Leu Ala Thr
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Cys Asn Gln Cys Ala Pro Ser Val Ala Asn Cys Pro Met Cys
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<210> 37
<211> 46
<212> PRT
<213> Cydia pomonella
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Glu Lys Glu Pro Gln Val Glu Asp Ser Lys Leu Cys Lys Ile Cys Tyr
Val Glu Glu Cys Ile Val Cys Phe Val Pro Cys Gly His Val Val Ala
Cys Ala Lys Cys Ala Leu Ser Val Asp Lys Cys Pro Met Cys
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<210> 38
<211> 46
<212> PRT
<213> Orgyia pseudotsugata
<400> 38
Ala Val Glu Ala Glu Val Ala Asp Asp Arg Leu Cys Lys Ile Cys Leu
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Cys Gly Lys Cys Ala Ala Gly Val Thr Thr Cys Pro Val Cys
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<212> DNA
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gacagegeet ttetageeaa getgatgaag agtgetgaea eetttgagtt gaagtatgae 240
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<211> 602

<212> PRT

<213> Mus musculus

<400> 40

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Ser Pro Met Glu Lys His Arg Lys Leu Tyr Pro Ser Cys Asn Phe Val Gln Thr Leu Asn Pro Ala Asn Ser Leu Glu Ala Ser Pro Arg Pro Ser 100 105 110 Leu Pro Ser Thr Ala Met Ser Thr Met Pro Leu Ser Phe Ala Ser Ser 120 125 Glu Asn Thr Gly Tyr Phe Ser Gly Ser Tyr Ser Ser Phe Pro Ser Asp 135 140 Pro Val Asn Phe Arg Ala Asn Gln Asp Cys Pro Ala Leu Ser Thr Ser 150 155 Pro Tyr His Phe Ala Met Asn Thr Glu Lys Ala Arg Leu Leu Thr Tyr 165 170 Glu Thr Trp Pro Leu Ser Phe Leu Ser Pro Ala Lys Leu Ala Lys Ala 180 185 Gly Phe Tyr Tyr Ile Gly Pro Gly Asp Arg Val Ala Cys Phe Ala Cys 200 Asp Gly Lys Leu Ser Asn Trp Glu Arg Lys Asp Asp Ala Met Ser Glu 215 220 His Gln Arg His Phe Pro Ser Cys Pro Phe Leu Lys Asp Leu Gly Gln 230 235 225 Ser Ala Ser Arg Tyr Thr Val Ser Asn Leu Ser Met Gln Thr His Ala 245 250 Ala Arg Ile Arg Thr Phe Ser Asn Trp Pro Ser Ser Ala Leu Val His 265 260 Ser Gln Glu Leu Ala Ser Ala Gly Phe Tyr Tyr Thr Gly His Ser Asp 280 Asp Val Lys Cys Leu Cys Cys Asp Gly Gly Leu Arg Cys Trp Glu Ser 295 300 Gly Asp Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg Cys Glu 310 315 Tyr Leu Leu Arg Ile Lys Gly Gln Glu Phe Val Ser Gln Val Gln Ala 325 330 Gly Tyr Pro His Leu Leu Glu Gln Leu Leu Ser Thr Ser Asp Ser Pro 345 Glu Asp Glu Asn Ala Asp Ala Ala Ile Val His Phe Gly Pro Gly Glu 360 Ser Ser Glu Asp Val Val Met Met Ser Thr Pro Val Val Lys Ala Ala 375 Leu Glu Met Gly Phe Ser Arg Ser Leu Val Arg Gln Thr Val Gln Trp 390 395 Gln Ile Leu Ala Thr Gly Glu Asn Tyr Arg Thr Val Ser Asp Leu Val Ile Gly Leu Leu Asp Ala Glu Asp Glu Met Arg Glu Glu Gln Met Glu 425 Gln Ala Ala Glu Glu Glu Ser Asp Asp Leu Ala Leu Ile Arg Lys Asn Lys Met Val Leu Phe Gln His Leu Thr Cys Val Thr Pro Met Leu 455 460 Tyr Cys Leu Leu Ser Ala Arg Ala Ile Thr Glu Gln Glu Cys Asn Ala Val Lys Gln Lys Pro His Thr Leu Gln Ala Ser Thr Leu Ile Asp Thr 490 Val Leu Ala Lys Gly Asn Thr Ala Ala Thr Ser Phe Arg Asn Ser Leu 505 500

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Arg Glu Ile Asp Pro Ala Leu Tyr Arg Asp Ile Phe Val Gln Gln Asp
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Ile Arg Ser Leu Pro Thr Asp Asp Ile Ala Ala Leu Pro Met Glu Glu
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                                             540
Gln Leu Arg Pro Leu Pro Glu Asp Arg Met Cys Lys Val Cys Met Asp
545
                    550
                                         555
                                                              560
Arg Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val Cys
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Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys Arg Gly Thr
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<210> 41 <211> 2416 <212> DNA <213> Mus musculus

<400> 41

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gaagggtagc attgtatatt taagcttagt ctgttgcaag ggaaggtcta tgctgttgag 2100 ctacaggact gtgtctgttc cagagcagga gttgggatgc ttgctgtatg tccttcagga 2160 cttcttggga tttgggaatt tggggaaagc tttggaatcc agtgatgtgg agctcagaaa 2220 tectggaace agtgaetetg gtaeteagta gatagggtae cetgtaette ttggtgettt 2280 tccagtctgg gaaataagga ggaatctgct gctggtaaaa atttgctgga tgtgagaaat 2340 agatgaaagt gtttcgggtg ggggcgtgca tcagtgtagt gtgtgcaggg atgtatgcag 2400 gccaaacact gtgtag

<210> 42

<211> 591

<212> PRT

<213> Mus musculus

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Gln Ala Arg Tyr Pro His Leu Leu Glu Gln Leu Leu Ser Thr Ser Asp
                                     330
Thr Pro Gly Glu Glu Asn Ala Asp Pro Thr Glu Thr Val Val His Phe
                                345
            340
                                                     350
Gly Pro Gly Glu Ser Ser Lys Asp Val Val Met Met Ser Thr Pro Val
                            360
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Val Lys Ala Ala Leu Glu Met Gly Phe Ser Arg Ser Leu Val Arg Gln
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Thr Val Gln Arg Gln Ile Leu Ala Thr Gly Glu Asn Tyr Arg Thr Val
                    390
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Asn Asp Ile Val Ser Val Leu Leu Asn Ala Glu Asp Glu Arg Arg Glu
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                                    410
Glu Glu Lys Glu Arg Gln Thr Glu Glu Met Ala Ser Gly Asp Leu Ser
                                425
                                                     430
            420
Leu Ile Arg Lys Asn Arg Met Ala Leu Phe Gln Gln Leu Thr His Val
                            440
                                                 445
Leu Pro Ile Leu Asp Asn Leu Leu Glu Ala Ser Val Ile Thr Lys Gln
                        455
Glu His Asp Ile Ile Arg Gln Lys Thr Gln Ile Pro Leu Gln Ala Arg
                    470
                                         475
Glu Leu Ile Asp Thr Val Leu Val Lys Gly Asn Ala Ala Asn Ile
                485
                                    490
Phe Lys Asn Ser Leu Lys Gly Ile Asp Ser Thr Leu Tyr Glu Asn Leu
            500
                                505
Phe Val Glu Lys Asn Met Lys Tyr Ile Pro Thr Glu Asp Val Ser Gly
                            520
Leu Ser Leu Glu Glu Gln Leu Arg Arg Leu Gln Glu Glu Arg Thr Cys
                        535
                                             540
Lys Val Cys Met Asp Arg Glu Val Ser Ile Val Phe Ile Pro Cys Gly
                    550
                                        555
His Leu Val Val Cys Gln Glu Cys Ala Pro Ser Leu Arg Lys Cys Pro
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Ile Cys Arg Gly Thr Ile Lys Gly Thr Val Arg Thr Phe Leu Ser
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<211> 11
<212> PRT
<213> artificial sequence based on Homo sapiens
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<400> 43

Met Glu Gln Lys Leu Ile Ser Glu Glu Asp Leu
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<210> 44

<211> 635

<212> PRT

<213> artificial sequence based on Homo sapiens, Mus musculus, Cydia pomonella, and Drosophila melanogaster

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<223> any amino acid or may be absent
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<222> (1)...(635)
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Leu Xaa Thr Phe Xaa Xaa Phe Pro Xaa Xaa Xaa Pro Val Ser Xaa Xaa
                        40
Xaa Leu Ala Arg Ala Gly Phe Xaa Tyr Thr Gly Xaa Xaa Asp Xaa Val
                    55
Xaa Cys Phe Xaa Cys Xaa Xaa Xaa Xaa Asp Xaa Trp Xaa Xaa Gly Asp
                 70
                                  75
Ser Xaa Xaa Xaa His Xaa Xaa Xaa Pro Xaa Cys Xaa Phe Ile
100
                           105
120
Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Tyr Xaa Xaa Xaa Xaa
                                     140
                    135
Xaa Xaa Xaa Xaa Arg Xaa Xaa Glu Xaa Xaa Xaa Xaa Xaa Xaa Xaa
                 150
                                  155
Xaa Xaa Xaa Xaa Xaa Asp Xaa Ser Asp Xaa Xaa Xaa Xaa Xaa Xaa
             165
                              170
Xaa Xaa Xaa Met Xaa Xaa Glu Glu Ala Arg Leu Xaa Thr Phe Xaa Xaa
          180
                           185
Trp Pro Xaa Xaa Xaa Xaa Leu Xaa Pro Xaa Glu Leu Ala Xaa Ala Gly
      195
                       200
Phe Tyr Tyr Xaa Gly Xaa Xaa Asp Xaa Val Xaa Cys Phe Xaa Cys Gly
                    215
                                     220
Gly Lys Leu Xaa Asn Trp Glu Pro Xaa Asp Xaa Ala Xaa Ser Glu His
                 230
                                  235
Xaa Arg His Phe Pro Xaa Cys Pro Phe Val Xaa Xaa Xaa Xaa Xaa Xaa
             245
                              250
260
                           265
                                            270
Ser Xaa Xaa Xaa Pro Xaa Asn Pro Xaa Met Ala Xaa Xaa Ala Arg
                       280
                                         285
Xaa Xaa Thr Phe Xaa Xaa Trp Pro Xaa Ser Xaa Xaa Val Xaa Xaa Glu
                    295
                                     300
Gln Leu Ala Xaa Ala Gly Phe Tyr Tyr Xaa Gly Xaa Gly Asp Xaa Val
                                  315
                 310
Lys Cys Phe Xaa Cys Xaa Gly Gly Leu Xaa Xaa Trp Xaa Xaa Xaa Asp
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                              330
Asp Pro Trp Xaa Gln His Ala Lys Trp Phe Pro Xaa Cys Xaa Tyr Leu
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Xaa Xaa Xaa Lys Gly Gln Glu Tyr Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa
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Xaa Xaa Leu Xaa Glu Xaa Leu Xaa Xaa Thr Xaa Xaa Xaa Xaa Xaa Xaa
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                            380
385
            390
                         395
Xaa Xaa Asp Xaa Val Xaa Xaa Xaa Pro Xaa Val Xaa Xaa Ala Xaa
          405
                      410
Xaa Met Gly Phe Xaa Xaa Xaa Xaa Val Lys Xaa Xaa Xaa Xaa Lys
                    425
Ile Xaa Xaa Xaa Gly Xaa Xaa Tyr Xaa Xaa Xaa Xaa Leu Val Xaa
     435
                 440
                              445
Asp Leu Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Glu Xaa Xaa Xaa
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470
500
                    505
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Xaa Xaa Xaa Gln Xaa Xaa Leu Gln Xaa Xaa Xaa Xaa Xaa Xaa
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535
                            540
555
570
          565
Gln Leu Arg Arg Leu Xaa Glu Glu Xaa Leu Cys Lys Xaa Cys Met Asp
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                    585
Xaa Glu Val Xaa Xaa Val Phe Xaa Pro Cys Gly His Leu Val Xaa Cys
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Xaa Xaa Cys Ala Xaa Ser Val Xaa Lys Cys Pro Met Cys Arg Xaa Xaa
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Ile Xaa Xaa Xaa Xaa Xaa Phe Leu Ser Xaa
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                         635
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<210> 45

<211> 204

<212> DNA

<213> Homo sapiens

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<210> 46

<211> 204

<212> DNA

<213> Homo sapiens

<213> Mus musculus

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<210> 47 <211> 198 <212> DNA <213> Homo	sapiens					
gcaagagctg	gattttatgc ctgattggaa	tttaqqtqaa	ggtgataaag	taaagtgctt	ggagcagctt tcactgtgga taaatggtat	120
<210> 48 <211> 138 <212> DNA <213> Homo	sapiens					
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<210> 49 <211> 204 <212> DNA <213> Mus 1	musculus					
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<210> 50 <211> 204 <212> DNA <213> Mus	musculus					
gagttagcta tgtgggggaa	atactaacct	ctactacaca ttgggaaccc	ggggctgatg	atcaagtgca	aacccccaga atgcttttgt acacaggaga	120
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<210> 52 <211> 138 <212> DNA <213> Mus musculus					
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<210> 53 <211> 204 <212> DNA <213> Homo sapiens					
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<210> 54 <211> 201 <212> DNA <213> Homo sapiens					
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<210> 55 <211> 204 <212> DNA <213> Homo sapiens					
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<210> 56 <211> 138 <212> DNA <213> Homo sapiens					

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<210> 57 <211> 203 <212> DNA					
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tgtgatggtg ggctgaggtg	ctgggaatct				180
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aggaagtgtc ccatctgt		_	-		138
<210> 61					
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<210> 65 <211> 204 <212> DNA <213> Mus	musculus					
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ggtgggaaac tgagcaactg ggaaccaaag gattatgcta tgtcagagca ccgcagacat 180
                                                                   201
tttccccact gtccatttct g
<210> 67
<211> 204
<212> DNA
<213> Mus musculus
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<211> 114
<212> DNA
<213> Mus musculus
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catctagtag tctgccagga atgtgcccct tctctaagga agtgccccat ctgc
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<211> 68
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<213> Homo sapiens
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Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu
                                                     30
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Gly Asp Thr Val Arg Cys Phe Ser Cys His Ala Ala Val Asp Arg Trp
                             40
Gln Tyr Gly Asp Ser Ala Val Gly Arg His Arg Lys Val Ser Pro Asn
                                             60
                        55
    50
Cys Arg Phe Ile
65
<210> 70
<211> 68
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<213> Homo sapiens
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Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr Ala His
Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr Gly Ile
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Gly Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp
Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn
                                             60
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Cys Phe Phe Val
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<211> 66
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Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp
Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro
Ser Glu Asp Pro Trp Glu Gln His Ala Lys Trp Tyr Pro Gly Cys Lys
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Tyr Leu
65
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213> Homo sapiens
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Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys Pro Met Cys
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                             40
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<213> Mus musculus
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Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu
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Gly Asp Thr Val Gln Cys Phe Ser Cys His Ala Ala Ile Asp Arg Trp
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                             40
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Cys Arg Phe Ile
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Asp Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp
Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn
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Cys Phe Phe Val
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Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp
Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro
Ser Glu Asp Pro Trp Asp Gln His Ala Lys Cys Tyr Pro Gly Cys Lys
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Tyr Leu
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Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val
Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp
                             40
Lys Arg Gly Asp Ser Pro Thr Glu Lys His Lys Lys Leu Tyr Pro Ser
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Cys Arg Phe Val
65
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Glu Asn Ala Arg Leu Leu Thr Phe Gln Thr Trp Pro Leu Thr Phe Leu
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Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu
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Pro Lys Asp Asn Ala Met Ser Glu His Leu Arg His Phe Pro Lys Cys
Pro Phe Ile
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<212> PRT
<213> Homo sapiens
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Val Asn Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Asn
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Ser Asp Asp Val Lys Cys Phe Cys Cys Asp Gly Gly Leu Arg Cys Trp
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Glu Ser Gly Asp Asp Pro Trp Val Gln His Ala Lys Trp Phe Pro Arg
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Cys Glu Tyr Leu
65
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<213> Homo sapiens
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Asp Lys Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val
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Cys Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys
                            40
<210> 81
<211> 68
<212> PRT
<213> Mus musculus
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Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Ala
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Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp
                             40
Lys Gln Gly Asp Ser Pro Met Glu Lys His Arg Lys Leu Tyr Pro Ser
                         55
Cys Asn Phe Val
<210> 82
<211> 67
<212> PRT
<213> Mus musculus
<400> 82
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Ser Pro Ala Lys Leu Ala Lys Ala Gly Phe Tyr Tyr Ile Gly Pro Gly
                                 25
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Asp Arg Val Ala Cys Phe Ala Cys Asp Gly Lys Leu Ser Asn Trp Glu
Arg Lys Asp Asp Ala Met Ser Glu His Gln Arg His Phe Pro Ser Cys
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Pro Phe Leu
65
<210> 83
<211> 68
<212> PRT
 <213> Mus musculus
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<400> 83
His Ala Ala Arg Ile Arg Thr Phe Ser Asn Trp Pro Ser Ser Ala Leu
Val His Ser Gln Glu Leu Ala Ser Ala Gly Phe Tyr Tyr Thr Gly His
Ser Asp Asp Val Lys Cys Leu Cys Cys Asp Gly Gly Leu Arg Cys Trp
Glu Ser Gly Asp Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg
                        55
Cys Glu Tyr Leu
65
<210> 84
<211> 46
<212> PRT
<213> Mus musculus
<400> 84
Glu Gln Leu Arg Pro Leu Pro Glu Asp Arg Met Cys Lys Val Cys Met
Asp Arg Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val
                                25
Cys Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys
<210> 85
<211> 68
<212> PRT
<213> Homo sapiens
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Glu Leu Tyr Arg Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro
Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val
Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp
                            40
Lys Leu Gly Asp Ser Pro Ile Gln Lys His Lys Gln Leu Tyr Pro Ser
                        55
Cys Ser Phe Ile
65
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<213> Homo sapiens
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Ser Pro Ser Glu Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly
                                25
Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu
                            40
Pro Lys Asp Asp Ala Met Ser Glu His Arg Arg His Phe Pro Asn Cys
    50
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Pro Phe Leu
65
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<211> 68
<212> PRT
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Val Gln Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Arg
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Asn Asp Asp Val Lys Cys Phe Gly Cys Asp Gly Gly Leu Arg Cys Trp
Glu Ser Gly Asp Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg
Cys Glu Phe Leu
65
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<212> PRT
<213> Homo sapiens
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Asp Lys Glu Val Ser Val Val Phe Ile Pro Cys Gly His Leu Val Val
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Cys Gln Glu Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys
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<211> 68

<212> PRT

<213> Mus musculus

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                            40
Lys Gln Gly Asp Ser Pro Val Glu Lys His Arg Gln Phe Tyr Pro Ser
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Cys Ser Phe Val
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Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu
        35
Pro Lys Asp Tyr Ala Met Ser Glu His Arg Arg His Phe Pro His Cys
                                             60
                         55
    50
Pro Phe Leu
65
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                             40
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                         55
    50
Cys Glu Phe Leu
65
<210> 92
<211> 38
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<213> Mus musculus
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### SEQUENCE LISTING

### (1) GENERAL INFORMATION:

- (i) APPLICANT: Korneluk, Robert G.
  Mackenzie, Alexander E.
  Baird, Stephen
- (ii) TITLE OF INVENTION: MAMMALIAN IAP GENE FAMILY, PRIMERS, PROBES, AND DETECTION METHODS
- (iii) NUMBER OF SEQUENCES: 42
- (iv) CORRESPONDENCE ADDRESS:
  - (A) ADDRESSEE: Fish & Richardson P.C.
  - (B) STREET: 225 Franklin Street
  - (C) CITY: Boston
  - (D) STATE: MA
  - (E) COUNTRY: USA
  - (F) ZIP: 02110-2804
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B) COMPUTER: IBM PC compatible
  - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
  - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER:
  - (B) FILING DATE:
  - (C) CLASSIFICATION:
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: US 08/511,485
  - (B) FILING DATE: 04-AUG-1995
  - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Clark, Paul T.
  - (B) REGISTRATION NUMBER: 30,162
  - (C) REFERENCE/DOCKET NUMBER: 07891/002001
  - (ix) TELECOMMUNICATION INFORMATION:
    - (A) TELEPHONE: 617/542-5070
    - (B) TELEFAX: 617/542-8906
    - (C) TELEX: 200154
- (2) INFORMATION FOR SEQ ID NO:1:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 46 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (ix) FEATURE:
    - (D) OTHER INFORMATION: Xaa at positions 2, 3, 4, 5,

6, 7, 9, 10, 11, 17, 18, 19, 20, 21, 23, 25, 30, 31, 32, 34, 35, 38, 39, 40, 41, 42, and 45 may be any amino acid. Xaa at position 8 is Glu or Asp. Xaa at positions 14 & 22 is Val or Ile.

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

Glu Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Xaa Cys Lys Xaa Cys Met
1 5 10 15

Xaa Xaa Xaa Xaa Xaa Xaa Phe Xaa Pro Cys Gly His Xaa Xaa Xaa 20 25 30

Cys Xaa Xaa Cys Ala Xaa Xaa Xaa Xaa Cys Pro Xaa Cys 35 40 45

#### (2) INFORMATION FOR SEQ ID NO:2:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 68 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (ix) FEATURE:
- (D) OTHER INFORMATION: Xaa at positions 1, 2, 3, 6, 9, 10, 14, 15, 18, 19, 20, 21, 24, 30, 32, 33, 35, 37, 40, 42, 43, 44, 45, 46, 47, 49, 50, 51, 53, 54, 55, 56, 57, 59, 60, 61, 62, 64 and 66 may be any amino acid. Xaa at positions 13, 16 and 17 may be any amino acid or may be absent.
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Xaa Xaa Xaa Arg Leu Xaa Thr Phe Xaa Xaa Trp Pro Xaa Xaa Xaa Xaa 1 1 15

Xaa Xaa Xaa Xaa Leu Ala Xaa Ala Gly Phe Tyr Tyr Xaa Gly Xaa 20 25 30

Xaa Asp Xaa Val Xaa Cys Phe Xaa Cys Xaa Xaa Xaa Xaa Xaa Xaa Trp 35 40 45

Cys Xaa Phe Val

## (2) INFORMATION FOR SEQ ID NO:3:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2540 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: DNA (genomic)

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

GAAAAGGTGG ACAAGTCCTA	TTTTCAAGAG	AAGATGACTT	TTAACAGTTT	TGAAGGATCT	60
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GCAGGGTTTC TTTATACTGG	TGAAGGAGAT	ACCGTGCGGT	GCTTTAGTTG	TCATGCAGCT	240
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ATCCAGAATG GTCAGTACAA	AGTTGAAAAC	TATCTGGGAA	GCAGAGATCA	TTTTGCCTTA	420
GACAGGCCAT CTGAGACACA	TGCAGACTAT	CTTTTGAGAA	CTGGGCAGGT	TGTAGATATA	480
TCAGACACCA TATACCCGAG	GAACCCTGCC	ATGTATTGTG	AAGAAGCTAG	ATTAAAGTCC	540
TTTCAGAACT GGCCAGACTA	TGCTCACCTA	ACCCCAAGAG	AGTTAGCAAG	TGCTGGACTC	60 <b>0</b>
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GTTTTGGGCC GGAATCTTAA	TATTCGAAGT	GAATCTGATG	CTGTGAGTTC	TGATAGGAAT	780
TTCCCAAATT CAACAAATCT	TCCAAGAAAT	CCATCCATGG	CAGATTATGA	AGCACGGATC	840
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TATGCTTTAG GTGAAGGTGA	TAAAGTAAAG	TGCTTTCACT	GTGGAGGAGG	GCTAACTGAT	960
TGGAAGCCCA GTGAAGACCC	TTGGGAACAA	CATGCTAAAT	GGTATCCAGG	GTGCAAATAT	1020
CTGTTAGAAC AGAAGGGACA	AGAATATATA	AACAATATTC	ATTTAACTCA	TTCACTTGAG	1080
GAGTGTCTGG TAAGAACTAC	TGAGAAAACA	CCATCACTAA	CTAGAAGAAT	TGATGATACC	1140
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TCATTACAGA AAGAGATTAG	TACTGAAGAG	CAGCTAAGGC	GCCTGCAAGA	GGAGAAGCTT	1380
TGCAAAATCT GTATGGATAG	AAATATTGCT	ATCGTTTTTG	TTCCTTGTGG	ACATCTAGTC	1440
ACTTGTAAAC AATGTGCTGA	AGCAGTTGAC	AAGTGTCCCA	TGTGCTACAC	AGTCATTACT	1500
TTCAAGCAAA AAATTTTTAT	GTCTTAATCT	AACTCTATAG	TAGGCATGTT	ATGTTGTTCT	1560
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TTTAATTGAA	ACCATAGACT	AAGAATAAGA	AGCATCATAC	TATAACTGAA	CACAATGTGT	1800
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TCTTTTCAGA	TAGGCTTAAC	AAATGGAGCT	TTCTGTATAT	AAATGTGGAG	ATTAGAGTTA	1920
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GAAAGATAGA	GATTGTTTTT	AGAGGTTGGT	TGTTGTGTTT	TAGGATTCTG	TCCATTTTCT	2040
TGTAAAGGGA	TAAACACGGA	CGTGTGCGAA	ATATGTTTGT	AAAGTGATTT	GCCATTGTTG	2100
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GAGATATGTT	AAGTGTAAAA	TGCAAGTGGC	GGGACACTAT	GTATAGTCTG	AGCCAGATCA	2220
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TTAAATGTGG	TTTCTCTTCG	GGGAGGGGG	GATTGGGGGA	GGGGCCCCAG	AGGGGTTTTA	2340
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## (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 497 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: unknown
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Thr Phe Asn Ser Phe Glu Gly Ser Lys Thr Cys Val Pro Ala Asp
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Ile Asn Lys Glu Glu Glu Phe Val Glu Glu Phe Asn Arg Leu Lys Thr 20 25 30

Phe Ala Asn Phe Pro Ser Gly Ser Pro Val Ser Ala Ser Thr Leu Ala 35 40 45

Arg Ala Gly Phe Leu Tyr Thr Gly Glu Gly Asp Thr Val Arg Cys Phe 50 55 60

Ser Cys His Ala Ala Val Asp Arg Trp Gln Tyr Gly Asp Ser Ala Val 65 70 75 80

Gly Arg His Arg Lys Val Ser Pro Asn Cys Arg Phe Ile Asn Gly Phe 85 90 95

Tyr Leu Glu Asn Ser Ala Thr Gln Ser Thr Asn Ser Gly Ile Gln Asn

			100					105					110		
Gly	Gln	Tyr 115	Lys	Val	Glu	Asn	Tyr 120	Leu	Gly	Ser	Arg	Asp 125	His	Phe	Ala
Leu	Asp 130	Arg	Pro	Ser	Glu	Thr 135	His	Ala	Asp	Tyr	Leu 140	Leu	Arg	Thr	Gly
Gln 145	Val	Val	Asp	Ile	Ser 150	Asp	Thr	Ile	Tyr	Pro 155	Arg	Asn	Pro	Ala	Met 160
Tyr	Сув	Glu	Glu	Ala 165	Arg	Leu	Lys	Ser	Phe 170	Gln	Asn	Trp	Pro	Asp 175	Tyr
Ala	His	Leu	Thr 180	Pro	Arg	Glu	Leu	Ala 185	Ser	Ala	Gly	Leu	Tyr 190	Tyr	Thr
Gly	Ile	Gly 195	Asp	Gln	Val	Gln	Cys 200	Phe	Cys	Cys	Gly	Gly 205	Lys	Leu	Lys
Asn	Trp 210	Glu	Pro	Cys	Asp	Arg 215	Ala	Trp	Ser	Glu	His 220	Arg	Arg	His	Phe
Pro 225	Asn	Cys	Phe	Phe	Val 230	Leu	Gly	Arg	Asn	Leu 235	Asn	Ile	Arg	Ser	Glu 240
Ser	Asp	Ala	Val	Ser 245	Ser	Asp	Arg	Asn	Phe 250	Pro	Asn	Ser	Thr	Asn 255	Leu
Pro	Arg	Asn	Pro 260	Ser	Met	Ala	Asp	Tyr 265	Glu	Ala	Arg	Ile	Phe 270	Thr	Phe
Gly	Thr	Trp 275	Ile	Tyr	Ser	Val	Asn 280	Lys	Glu	Gln	Leu	Ala 285	Arg	Ala	Gly
Phe	Tyr 290	Ala	Leu	Gly	Glu	Gly 295	Asp	Lys	Val	Lys	Cys 300	Phe	His	Cys	Gly
Gly 305	Gly	Leu	Thr	Asp	Trp 310	Lys	Pro	Ser	Glu	Asp 315	Pro	Trp	Glu	Gln	His 320
Ala	Lys	Trp	Tyr	Pro 325	Gly	Cys	Lys	Tyr	Leu 330	Leu	Glu	Gln	Lys	Gly 335	Gln
Glu	Tyr	Ile	Asn 340	Asn	Ile	His	Leu	Thr 345	His	Ser	Leu	Glu	Glu 350	Cys	Leu
Val	Arg	Thr 355	Thr	Glu	Lys	Thr	Pro 360	Ser	Leu	Thr	Arg	Arg 365	Ile	Asp	Asp
Thr	Ile 370	Phe	Gln	Asn	Pro	Met 375	Val	Gln	Glu	Ala	Ile 380	Arg	Met	Gly	Phe
Ser 385	Phe	Lys	Asp	Ile	Lys 390	Lys	Ile	Met	Glu	Glu 395	Lys	Ile	Gln	Ile	Ser 400
Gly	Ser	Asn	Tyr	Lys 405	Ser	Leu	Glu	Val	Leu 410	Val	Ala	Asp	Leu	Val 415	Asn
Ala	Gln	Lys	Asp 420	Ser	Met	Gln	Asp	Glu 425	Ser	Ser	Gln	Thr	Ser 430	Leu	Gln

Lys Glu Ile Ser Thr Glu Glu Gln Leu Arg Arg Leu Gln Glu Glu Lys 435 440 445

Leu Cys Lys Ile Cys Met Asp Arg Asn Ile Ala Ile Val Phe Val Pro 450 455 460

Cys Gly His Leu Val Thr Cys Lys Gln Cys Ala Glu Ala Val Asp Lys 465 470 475 480

Cys Pro Met Cys Tyr Thr Val Ile Thr Phe Lys Gln Lys Ile Phe Met 485 490 495

Ser

### (2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2676 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: DNA (genomic)

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

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ATTTCATTAT	GAACATAGTA	GAAAACAGCA	TATTCTTATC	AAATTTGATG	AAAAGCGCCA	240
ACACGTTTGA	ACTGAAATAC	GACTTGTCAT	GTGAACTGTA	CCGAATGTCT	ACGTATTCCA	300
CTTTTCCTGC	TGGGGTTCCT	GTCTCAGAAA	GGAGTCTTGC	TCGTGCTGGT	TTCTATTACA	360
CTGGTGTGAA	TGACAAGGTC	AAATGCTTCT	GTTGTGGCCT	GATGCTGGAT	AACTGGAAAA	420
GAGGAGACAG	TCCTACTGAA	AAGCATAAAA	AGTTGTATCC	TAGCTGCAGA	TTCGTTCAGA	480
GTCTAAATTC	CGTTAACAAC	TTGGAAGCTA	CCTCTCAGCC	TACTTTTCCT	TCTTCAGTAA	540
CACATTCCAC	ACACTCATTA	CTTCCGGGTA	CAGAAAACAG	TGGATATTTC	CGTGGCTCTT	600
ATTCAAACTC	TCCATCAAAT	CCTGTAAACT	CCAGAGCAAA	TCAAGAATTT	TCTGCCTTGA	660
TGAGAAGTTC	CTACCCCTGT	CCAATGAATA	ACGAAAATGC	CAGATTACTT	ACTTTTCAGA	720
CATGGCCATT	GACTTTTCTG	TCGCCAACAG	ATCTGGCACG	AGCAGGCTTT	TACTACATAG	780
GACCTGGAGA	CAGAGTGGCT	TGCTTTGCCT	GTGGTGGAAA	ATTGAGCAAT	TGGGAACCGA	840
AGGATAATGC	TATGTCAGAA	CACCTGAGAC	ATTTTCCCAA	ATGCCCATTT	ATAGAAAATC	900
AGCTTCAAGA	CACTTCAAGA	TACACAGTTT	CTAATCTGAG	CATGCAGACA	CATGCAGCCC	960

GACTCAGGTG TTGGGAATCT GGAGATGATC CATGGGTTCA ACATGCCAAG TGGTTTCCAA  GGTGTGAGTA CTTGATAAGA ATTAAAGGAC AGGAGTTCAT CCGTCAAGTT CAAGCCAGTT  ACCCTCATCT ACTTGAACAG CTGCTATCCA CATCAGACAG CCCAGGAGAT GAAAATGCAG  AGTCATCAAT TATCCATTTG GAACCTGGAG AAGACCATTC AGAAGATGCA ATCATGATGA  1	1080 1140 1200 1260 1320 1380 1440
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	.380 .440
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ATGTTTCAGA TCTACCAGTG GAAGAACAAT TGCGGAGACT ACCAGAAGAA AGAACATGTA 1	.860
AAGTGTGTAT GGACAAGAA GTGTCCATAG TGTTTATTCC TTGTGGTCAT CTAGTAGTAT 1	920
GCAAAGATTG TGCTCCTTCT TTAAGAAAGT GTCCTATTTG TAGGAGTACA ATCAAGGGTA 1	980
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TTATTAAATG TATTATAACT TTAACTTTTA TCCTAATTTG GTTTCCTTAA AATTTTTATT 2	100
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CAGTGTCCTA TACATCGAAG GTGTGCATAT ATGTTGAATC ACATTTTAGG GACATGGTGT 2	580
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# (2) INFORMATION FOR SEQ ID NO:6:

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 604 amino acids

- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:
- Met Asn Ile Val Glu Asn Ser Ile Phe Leu Ser Asn Leu Met Lys Ser 1 10 15
- Ala Asn Thr Phe Glu Leu Lys Tyr Asp Leu Ser Cys Glu Leu Tyr Arg 20 25 30
- Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro Val Ser Glu Arg 35 40 45
- Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val Asn Asp Lys Val 50 55 60
- Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp Lys Arg Gly Asp 65 70 75 80
- Ser Pro Thr Glu Lys His Lys Leu Tyr Pro Ser Cys Arg Phe Val 85 90 95
- Gln Ser Leu Asn Ser Val Asn Asn Leu Glu Ala Thr Ser Gln Pro Thr 100 105 110
- Phe Pro Ser Ser Val Thr His Ser Thr His Ser Leu Leu Pro Gly Thr 115 120 125
- Glu Asn Ser Gly Tyr Phe Arg Gly Ser Tyr Ser Asn Ser Pro Ser Asn 130 135 140
- Pro Val Asn Ser Arg Ala Asn Gln Glu Phe Ser Ala Leu Met Arg Ser 145 150 155 160
- Ser Tyr Pro Cys Pro Met Asn Asn Glu Asn Ala Arg Leu Leu Thr Phe 165 170 175
- Gln Thr Trp Pro Leu Thr Phe Leu Ser Pro Thr Asp Leu Ala Arg Ala 180 185 190
- Gly Phe Tyr Tyr Ile Gly Pro Gly Asp Arg Val Ala Cys Phe Ala Cys 195 200 205
- Gly Gly Lys Leu Ser Asn Trp Glu Pro Lys Asp Asn Ala Met Ser Glu 210 215 220
- His Leu Arg His Phe Pro Lys Cys Pro Phe Ile Glu Asn Gln Leu Gln 225 230 235 240
- Asp Thr Ser Arg Tyr Thr Val Ser Asn Leu Ser Met Gln Thr His Ala 245 250 255
- Ala Arg Phe Lys Thr Phe Phe Asn Trp Pro Ser Ser Val Leu Val Asn 260 265 270

Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Asn Ser Asp Asp Val Lys Cys Phe Cys Cys Asp Gly Gly Leu Arg Cys Trp Glu Ser Gly Asp Asp Pro Trp Val Gln His Ala Lys Trp Phe Pro Arg Cys Glu Tyr Leu Ile Arg Ile Lys Gly Gln Glu Phe Ile Arg Gln Val Gln Ala Ser Tyr Pro His Leu Leu Glu Gln Leu Leu Ser Thr Ser Asp Ser Pro 345 Gly Asp Glu Asn Ala Glu Ser Ser Ile Ile His Leu Glu Pro Gly Glu 360 Asp His Ser Glu Asp Ala Ile Met Met Asn Thr Pro Val Ile Asn Ala 375 Ala Val Glu Met Gly Phe Ser Arg Ser Leu Val Lys Gln Thr Val Gln 390 Arg Lys Ile Leu Ala Thr Gly Glu Asn Tyr Arg Leu Val Asn Asp Leu Val Leu Asp Leu Leu Asn Ala Glu Asp Glu Ile Arg Glu Glu Glu Arg Glu Arg Ala Thr Glu Glu Lys Glu Ser Asn Asp Leu Leu Leu Ile Arg Lys Asn Arg Met Ala Leu Phe Gln His Leu Thr Cys Val Ile Pro Ile 455 Leu Asp Ser Leu Leu Thr Ala Gly Ile Ile Asn Glu Gln Glu His Asp Val Ile Lys Gln Lys Thr Gln Thr Ser Leu Gln Ala Arg Glu Leu Ile 485 490 Asp Thr Ile Leu Val Lys Gly Asn Ile Ala Ala Thr Val Phe Arg Asn 505 Ser Leu Gln Glu Ala Glu Ala Val Leu Tyr Glu His Leu Phe Val Gln 515 Gln Asp Ile Lys Tyr Ile Pro Thr Glu Asp Val Ser Asp Leu Pro Val Glu Glu Gln Leu Arg Arg Leu Pro Glu Glu Arg Thr Cys Lys Val Cys 550 Met Asp Lys Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val Cys Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys Arg Ser Thr Ile Lys Gly Thr Val Arg Thr Phe Leu Ser

## (2) INFORMATION FOR SEQ ID NO:7:

- (i) SEQUENCE CHARACTERISTICS:
  - $(\bar{A})$  LENGTH: 2580 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: DNA (genomic)

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

TTAGGTTACC	TGAAAGAGTT	ACTACAACCC	CAAAGAGTTG	TGTTCTAAGT	AGTATCTTGG	60
TAATTCAGAG	AGATACTCAT	CCTACCTGAA	TATAAACTGA	GATAAATCCA	GTAAAGAAAG	120
TGTAGTAAAT	TCTACATAAG	AGTCTATCAT	TGATTTCTTT	TTGTGGTGGA	AATCTTAGTT	180
CATGTGAAGA	AATTTCATGT	GAATGTTTTA	GCTATCAAAC	AGTACTGTCA	CCTACTCATG	240
CACAAAACTG	CCTCCCAAAG	ACTTTTCCCA	GGTCCCTCGT	ATCAAAACAT	TAAGAGTATA	300
ATGGAAGATA	GCACGATCTT	GTCAGATTGG	ACAAACAGCA	ACAAACAAAA	AATGAAGTAT	360
GACTTTTCCT	GTGAACTCTA	CAGAATGTCT	ACATATTCAA	CTTTCCCCGC	CGGGGTGCCT	420
GTCTCAGAAA	GGAGTCTTGC	TCGTGCTGGT	TTTTATTATA	CTGGTGTGAA	TGACAAGGTC	480
AAATGCTTCT	GTTGTGGCCT	GATGCTGGAT	AACTGGAAAC	TAGGAGACAG	TCCTATTCAA	540
AAGCATAAAC	AGCTATATCC	TAGCTGTAGC	TTTATTCAGA	ATCTGGTTTC	AGCTAGTCTG	600
GGATCCACCT	CTAAGAATAC	GTCTCCAATG	AGAAACAGTT	TTGCACATTC	ATTATCTCCC	660
ACCTTGGAAC	ATAGTAGCTT	GTTCAGTGGT	TCTTACTCCA	GCCTTCCTCC	AAACCCTCTT	720
AATTCTAGAG	CAGTTGAAGA	CATCTCTTCA	TCGAGGACTA	ACCCCTACAG	TTATGCAATG	780
AGTACTGAAG	AAGCCAGATT	TCTTACCTAC	CATATGTGGC	CATTAACTTT	TTTGTCACCA	840
TCAGAATTGG	CAAGAGCTGG	TTTTTATTAT	ATAGGACCTG	GAGATAGGGT	AGCCTGCTTT	900
GCCTGTGGTG	GGAAGCTCAG	TAACTGGGAA	CCAAAGGATG	ATGCTATGTC	AGAACACCGG	960
AGGCATTTTC	CCAACTGTCC	ATTTTTGGAA	AATTCTCTAG	AAACTCTGAG	GTTTAGCATT	1020
TCAAATCTGA	GCATGCAGAC	ACATGCAGCT	CGAATGAGAA	CATTTATGTA	CTGGCCATCT	1080
AGTGTTCCAG	TTCAGCCTGA	GCAGCTTGCA	AGTGCTGGTT	TTTATTATGT	GGGTCGCAAT	1140
GATGATGTCA	AATGCTTTGG	TTGTGATGGT	GGCTTGAGGT	GTTGGGAATC	TGGAGATGAT	1200
CCATGGGTAG	AACATGCCAA	GTGGTTTCCA	AGGTGTGAGT	TCTTGATACG	AATGAAAGGC	1260
CAAGAGTTTG	TTGATGAGAT	TCAAGGTAGA	TATCCTCATC	TTCTTGAACA	GCTGTTGTCA	1320
ACTTCAGATA	CCACTGGAGA	AGAAAATGCT	GACCCACCAA	TTATTCATTT	TGGACCTGGA	1380
GAAAGTTCTT	CAGAAGATGC	TGTCATGATG	AATACACCTG	TGGTTAAATC	TGCCTTGGAA	1440

ATGGGCTTTA ATAGAGACCT GGTGAAACAA ACAGTTCTAA GTAAAATCCT GACAACTGGA 1500 GAGAACTATA AAACAGTTAA TGATATTGTG TCAGCACTTC TTAATGCTGA AGATGAAAAA 1560 AGAGAAGAG AGAAGGAAAA ACAAGCTGAA GAAATGGCAT CAGATGATTT GTCATTAATT 1620 CGGAAGAACA GAATGGCTCT CTTTCAACAA TTGACATGTG TGCTTCCTAT CCTGGATAAT 1680 CTTTTAAAGG CCAATGTAAT TAATAAACAG GAACATGATA TTATTAAACA AAAAACACAG 1740 ATACCTTTAC AAGCGAGAGA ACTGATTGAT ACCATTTGGG TTAAAGGAAA TGCTGCGGCC 1800 AACATCTTCA AAAACTGTCT AAAAGAAATT GACTCTACAT TGTATAAGAA CTTATTTGTG 1860 GATAAGAATA TGAAGTATAT TCCAACAGAA GATGTTTCAG GTCTGTCACT GGAAGAACAA 1920 TTGAGGAGGT TGCAAGAAGA ACGAACTTGT AAAGTGTGTA TGGACAAAGA AGTTTCTGTT 1980 GTATTTATTC CTTGTGGTCA TCTGGTAGTA TGCCAGGAAT GTGCCCCTTC TCTAAGAAAA 2040 TGCCCTATTT GCAGGGGTAT AATCAAGGGT ACTGTTCGTA CATTTCTCTC TTAAAGAAAA 2100 ATAGTCTATA TTTTAACCTG CATAAAAAGG TCTTTAAAAT ATTGTTGAAC ACTTGAAGCC 2160 ATCTAAAGTA AAAAGGGAAT TATGAGTTTT TCAATTAGTA ACATTCATGT TCTAGTCTGC 2220 TTTGGTACTA ATAATCTTGT TTCTGAAAAG ATGGTATCAT ATATTTAATC TTAATCTGTT 2280 TATTTACAAG GGAAGATTTA TGTTTGGTGA ACTATATTAG TATGTATGTG TACCTAAGGG 2340 AGTAGCGTCN CTGCTTGTTA TGCATCATTT CAGGAGTTAC TGGATTTGTT GTTCTTTCAG 2400 AAAGCTTTGA ANACTAAATT ATAGTGTAGA AAAGAACTGG AAACCAGGAA CTCTGGAGTT 2460 CATCAGAGTT ATGGTGCCGA ATTGTCTTTG GTGCTTTTCA CTTGTGTTTT AAAATAAGGA 2520 TTTTTCTCTT ATTTCTCCCC CTAGTTTGTG AGAAACATCT CAATAAAGTG CTTTAAAAAG 2580

## (2) INFORMATION FOR SEQ ID NO:8:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 618 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Met His Lys Thr Ala Ser Gln Arg Leu Phe Pro Gly Pro Ser Tyr Gln 1 5 10 15

Asn Ile Lys Ser Ile Met Glu Asp Ser Thr Ile Leu Ser Asp Trp Thr 20 25 30

Asn Ser Asn Lys Gln Lys Met Lys Tyr Asp Phe Ser Cys Glu Leu Tyr 35 40 45

Arg Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro Val Ser Glu 50 60

Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val Asn Asp Lys 70 75 80

Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp Lys Leu Gly 85 90 95

Asp Ser Pro Ile Gln Lys His Lys Gln Leu Tyr Pro Ser Cys Ser Phe 100 105 110

Ile Gln Asn Leu Val Ser Ala Ser Leu Gly Ser Thr Ser Lys Asn Thr 115 120 125

Ser Pro Met Arg Asn Ser Phe Ala His Ser Leu Ser Pro Thr Leu Glu 130 135 140

His Ser Ser Leu Phe Ser Gly Ser Tyr Ser Ser Leu Pro Pro Asn Pro 145 150 155 160

Leu Asn Ser Arg Ala Val Glu Asp Ile Ser Ser Ser Arg Thr Asn Pro 165 170 175

Tyr Ser Tyr Ala Met Ser Thr Glu Glu Ala Arg Phe Leu Thr Tyr His 180 185 190

Met Trp Pro Leu Thr Phe Leu Ser Pro Ser Glu Leu Ala Arg Ala Gly
195 200 205

Phe Tyr Tyr Ile Gly Pro Gly Asp Arg Val Ala Cys Phe Ala Cys Gly 210 215 220

Gly Lys Leu Ser Asn Trp Glu Pro Lys Asp Asp Ala Met Ser Glu His 225 235 240

Arg Arg His Phe Pro Asn Cys Pro Phe Leu Glu Asn Ser Leu Glu Thr 245 250 255

Leu Arg Phe Ser Ile Ser Asn Leu Ser Met Gln Thr His Ala Ala Arg 260 265 270

Met Arg Thr Phe Met Tyr Trp Pro Ser Ser Val Pro Val Gln Pro Glu 275 280 285

Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Arg Asn Asp Asp Val 290 295 300

Lys Cys Phe Gly Cys Asp Gly Gly Leu Arg Cys Trp Glu Ser Gly Asp 305 310 315 320

Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg Cys Glu Phe Leu 325 330 335

Ile Arg Met Lys Gly Gln Glu Phe Val Asp Glu Ile Gln Gly Arg Tyr 340 345 350

Pro His Leu Leu Glu Gln Leu Leu Ser Thr Ser Asp Thr Thr Gly Glu 355 360 365

Glu Asn Ala Asp Pro Pro Ile Ile His Phe Gly Pro Gly Glu Ser Ser 370 380

Ser Glu Asp Ala Val Met Met Asn Thr Pro Val Val Lys Ser Ala Leu 385 Glu Met Gly Phe Asn Arg Asp Leu Val Lys Gln Thr Val Leu Ser Lys 405 Ile Leu Thr Thr Gly Glu Asn Tyr Lys Thr Val Asn Asp Ile Val Ser 420 Ala Leu Leu Asn Ala Glu Asp Glu Lys Arg Glu Glu Glu Lys Glu Lys 440 Gln Ala Glu Glu Met Ala Ser Asp Asp Leu Ser Leu Ile Arg Lys Asn 450 Arg Met Ala Leu Phe Gln Gln Leu Thr Cys Val Leu Pro Ile Leu Asp 470 Asn Leu Leu Lys Ala Asn Val Ile Asn Lys Gln Glu His Asp Ile Ile 495 490 Lys Gln Lys Thr Gln Ile Pro Leu Gln Ala Arg Glu Leu Ile Asp Thr 505 500 Ile Trp Val Lys Gly Asn Ala Ala Ala Asn Ile Phe Lys Asn Cys Leu 520 Lys Glu Ile Asp Ser Thr Leu Tyr Lys Asn Leu Phe Val Asp Lys Asn Met Lys Tyr Ile Pro Thr Glu Asp Val Ser Gly Leu Ser Leu Glu Glu 550 545 Gln Leu Arg Arg Leu Gln Glu Glu Arg Thr Cys Lys Val Cys Met Asp 570 Lys Glu Val Ser Val Val Phe Ile Pro Cys Gly His Leu Val Val Cys 585 580 Gln Glu Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys Arg Gly Ile 600 595 Ile Lys Gly Thr Val Arg Thr Phe Leu Ser 615

#### (2) INFORMATION FOR SEQ ID NO:9:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2100 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: both
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

  GACACTCTGC TGGGCGGGG GCCGCCCTCC TCCGGGACCT CCCCTCGGGA ACCGTCGCCC

GCGGCGCTTA GTTAGGACTG GAGTGCTTGG CGCGAAAAGG TGGACAAGTC CTATTTTCCA 120 GAGAAGATGA CTTTTAACAG TTTTGAAGGA ACTAGAACTT TTGTACTTGC AGACACCAAT 180 AAGGATGAAG AATTTGTAGA AGAGTTTAAT AGATTAAAAA CATTTGCTAA CTTCCCAAGT 240 AGTAGTCCTG TTTCAGCATC AACATTGGCG CGAGCTGGGT TTCTTTATAC CGGTGAAGGA 300 GACACCGTGC AATGTTTCAG TTGTCATGCG GCAATAGATA GATGGCAGTA TGGAGACTCA 360 GCTGTTGGAA GACACAGGAG AATATCCCCA AATTGCAGAT TTATCAATGG TTTTTATTTT 420 GAAAATGGTG CTGCACAGTC TACAAATCCT GGTATCCAAA ATGGCCAGTA CAAATCTGAA 480 AACTGTGTGG GAAATAGAAA TCCTTTTGCC CCTGACAGGC CACCTGAGAC TCATGCTGAT 540 TATCTCTTGA GAACTGGACA GGTTGTAGAT ATTTCAGACA CCATATACCC GAGGAACCCT 600 GCCATGTGTA GTGAAGAAGC CAGATTGAAG TCATTTCAGA ACTGGCCGGA CTATGCTCAT 660 TTAACCCCCA GAGAGTTAGC TAGTGCTGGC CTCTACTACA CAGGGGCTGA TGATCAAGTG 720 CAATGCTTTT GTTGTGGGGG AAAACTGAAA AATTGGGAAC CCTGTGATCG TGCCTGGTCA 780 GAACACAGGA GACACTTTCC CAATTGCTTT TTTGTTTTGG GCCGGAACGT TAATGTTCGA 840 900 AGTGAATCTG GTGTGAGTTC TGATAGGAAT TTCCCAAATT CAACAAACTC TCCAAGAAAT CCAGCCATGG CAGAATATGA AGCACGGATC GTTACTTTTG GAACATGGAT ATACTCAGTT 960 AACAAGGAGC AGCTTGCAAG AGCTGGATTT TATGCTTTAG GTGAAGGCGA TAAAGTGAAG 1020 TGCTTCCACT GTGGAGGAGG GCTCACGGAT TGGAAGCCCAA GTGAAGACCC CTGGGACCAG 1080 CATGCTAAGT GCTACCCAGG GTGCAAATAC CTATTGGATG AGAAGGGGCA AGAATATATA 1140 AATAATATTC ATTTAACCCA TCCACTTGAG GAATCTTTGG GAAGAACTGC TGAAAAAACA 1200 CCACCGCTAA CTAAAAAAAT CGATGATACC ATCTTCCAGA ATCCTATGGT GCAAGAAGCT 1260 ATACGAATGG GATTTAGCTT CAAGGACCTT AAGAAAACAA TGGAAGAAAA AATCCAAACA 1320 TCCGGGAGCA GCTATCTATC ACTTGAGGTC CTGATTGCAG ATCTTGTGAG TGCTCAGAAA 1380 GATAATACGG AGGATGAGTC AAGTCAAACT TCATTGCAGA AAGACATTAG TACTGAAGAG 1440 CAGCTAAGGC GCCTACAAGA GGAGAAGCTT TCCAAAATCT GTATGGATAG AAATATTGCT 1500 ATCGTTTTTT TTCCTTGTGG ACATCTGGCC ACTTGTAAAC AGTGTGCAGA AGCAGTTGAC 1560 AAATGTCCCA TGTGCTACAC CGTCATTACG TTCAACCAAA AAATTTTTAT GTCTTAGTGG 1620 GGCACCACAT GTTATGTTCT TCTTGCTCTA ATTGAATGTG TAATGGGAGC GAACTTTAAG 1680 TAATCCTGCA TTTGCATTCC ATTAGCATCC TGCTGTTTCC AAATGGAGAC CAATGCTAAC 1740 AGCACTGTTT CCGTCTAAAC ATTCAATTTC TGGATCTTTC GAGTTATCAG CTGTATCATT 1800 TAGCCAGTGT TTTACTCGAT TGAAACCTTA GACAGAGAAG CATTTTATAG CTTTTCACAT 1860 GTATATTGGT AGTACACTGA CTTGATTTCT ATATGTAAGT GAATTCATCA CCTGCATGTT 1920

TCATGCCTTT TGCATAAGCT TAACAAATGG AGTGTTCTGT ATAAGCATGG AGATGTGATG 1980
GAATCTGCCC AATGACTTTA ATTGGCTTAT TGTAAACACG GAAAGAACTG CCCCACGCTG 2040
CTGGGAGGAT AAAGATTGTT TTAGATGCTC ACTTCTGTGT TTTAGGATTC TGCCCATTTA 2100

#### (2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 496 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Met Thr Phe Asn Ser Phe Glu Gly Thr Arg Thr Phe Val Leu Ala Asp 1 5 10 15

Thr Asn Lys Asp Glu Glu Phe Val Glu Glu Phe Asn Arg Leu Lys Thr 20 25 30

Phe Ala Asn Phe Pro Ser Ser Ser Pro Val Ser Ala Ser Thr Leu Ala 35 40 45

Arg Ala Gly Phe Leu Tyr Thr Gly Glu Gly Asp Thr Val Gln Cys Phe 50 60

Ser Cys His Ala Ala Ile Asp Arg Trp Gln Tyr Gly Asp Ser Ala Val 65 70 75 80

Gly Arg His Arg Arg Ile Ser Pro Asn Cys Arg Phe Ile Asn Gly Phe 85 90 95

Tyr Phe Glu Asn Gly Ala Ala Gln Ser Thr Asn Pro Gly Ile Gln Asn 100 105 110

Gly Gln Tyr Lys Ser Glu Asn Cys Val Gly Asn Arg Asn Pro Phe Ala 115 120 125

Pro Asp Arg Pro Pro Glu Thr His Ala Asp Tyr Leu Leu Arg Thr Gly 130 140

Gln Val Val Asp Ile Ser Asp Thr Ile Tyr Pro Arg Asn Pro Ala Met 145 150 155 160

Cys Ser Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr 165 170 175

Ala His Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr 180 185 190

Gly Ala Asp Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys 195 200 205

Asn Trp Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe

	210					215					220				
Pro 225	Asn	Cys	Phe	Phe	Val 230	Leu	Gly	Arg	Asn	Val 235	Asn	Val	Arg	Ser	Glu 240
Ser	Gly	Val	Ser	Ser 245	Asp	Arg	Asn	Phe	Pro 250	Asn	Ser	Thr	Asn	Ser 255	Pro
Arg	Asn	Pro	Ala 260	Met	Ala	Glu	Tyr	Glu 265	Ala	Arg	Ile	Val	Thr 270	Phe	Gly
Thr	Trp	Ile 275	Tyr	Ser	Val	Asn	Lys 280	Glu	Gln	Leu	Ala	Arg 285	Ala	Gly	Phe
Tyr	Ala 290	Leu	Gly	Glu	Gly	Asp 295	Lys	Val	Lys	Cys	Phe 300	His	Cys	Gly	Gly
Gly 305	Leu	Thr	Asp	Trp	Lys 310	Pro	Ser	Glu	Asp	Pro 315	Trp	Asp	Gln	His	Ala 320
Lys	Cys	Tyr	Pro	Gly 325	Cys	Lys	Tyr	Leu	Leu 330	Asp	Glu	Lys	Gly	Gln 335	Glu
Tyr	Ile	Asn	Asn 340	Ile	His	Leu	Thr	His 345	Pro	Leu	Glu	Glu	Ser 350	Leu	Gly
Arg	Thr	Ala 355	Glu	Lys	Thr	Pro	Pro 360	Leu	Thr	Lys	Lys	Ile 365	Asp	Asp	Thr
Ile	Phe 370	Gln	Asn	Pro	Met	Val 375	Gln	Glu	Ala	Ile	Arg 380	Met	Gly	Phe	Ser
Phe 385	Lys	Asp	Leu	Lys	Lys 390	Thr	Met	Glu	Glu	Lys 395	Ile	Gln	Thr	Ser	Gly 400
Ser	Ser	Tyr	Leu	Ser 405	Leu	Glu	Val	Leu	Ile 410	Ala	Asp	Leu	Val	Ser 415	Ala
Gln	Lys	Asp	Asn 420	Thr	Glu	Asp	Glu	Ser 425	Ser	Gln	Thr	Ser	Leu 430	Gln	Lys
Asp	Ile	Ser 435	Thr	Glu	Glu	Gln	Leu 440	Arg	Arg	Leu	Gln	Glu 445	Glu	Lys	Leu
Ser	Lys 450	Ile	Cys	Met	Asp	Arg 455	Asn	Ile	Ala	Ile	Val 460	Phe	Phe	Pro	Сув
Gly 465	His	Leu	Ala	Thr	Cys 470	Lys	Gln	Cys	Ala	Glu 475	Ala	Val	Asp	Lys	Cys 480
Pro	Met	Cys	Tyr	Thr 485	Val	Ile	Thr	Phe	Asn 490	Gln	Lys	Ile	Phe	Met 495	Ser

## (2) INFORMATION FOR SEQ ID NO:11:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 67 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant

(D) TOPOLOGY: both

#### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

Lys Ala Ala Arg Leu Gly Thr Tyr Thr Asn Trp Pro Val Gln Phe Leu 10 15

Glu Pro Ser Arg Met Ala Ala Ser Gly Phe Tyr Tyr Leu Gly Arg Gly 20 25 30

Asp Glu Val Arg Cys Ala Phe Cys Lys Val Glu Ile Thr Asn Trp Val 35 40 45

Arg Gly Asp Asp Pro Glu Thr Asp His Lys Arg Trp Ala Pro Gln Cys 50 55 60

Pro Phe Val

## (2) INFORMATION FOR SEQ ID NO:12:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 275 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Met Ser Asp Leu Arg Leu Glu Glu Val Arg Leu Asn Thr Phe Glu Lys
1 10 15

Trp Pro Val Ser Phe Leu Ser Pro Glu Thr Met Ala Lys Asn Gly Phe 20 25 30

Tyr Tyr Leu Gly Arg Ser Asp Glu Val Arg Cys Ala Phe Cys Lys Val 35 40 45

Glu Ile Met Arg Trp Lys Glu Gly Glu Asp Pro Ala Ala Asp His Lys 50 55 60

Lys Trp Ala Pro Gln Cys Pro Phe Val Lys Gly Ile Asp Val Cys Gly 65 70 75 80

Ser Ile Val Thr Thr Asn Asn Ile Gln Asn Thr Thr Thr His Asp Thr 85 90 95

Ile Ile Gly Pro Ala His Pro Lys Tyr Ala His Glu Ala Ala Arg Val 100 105 110

Lys Ser Phe His Asn Trp Pro Arg Cys Met Lys Gln Arg Pro Glu Gln 115 120 125

Met Ala Asp Ala Gly Phe Phe Tyr Thr Gly Tyr Gly Asp Asn Thr Lys

140 135 130 Cys Phe Tyr Cys Asp Gly Gly Leu Lys Asp Trp Glu Pro Glu Asp Val 155 150 Pro Trp Glu Gln His Val Arg Trp Phe Asp Arg Cys Ala Tyr Val Gln 165 Leu Val Lys Gly Arg Asp Tyr Val Gln Lys Val Ile Thr Glu Ala Cys 185 Val Leu Pro Gly Glu Asn Thr Thr Val Ser Thr Ala Ala Pro Val Ser 200 Glu Pro Ile Pro Glu Thr Lys Ile Glu Lys Glu Pro Gln Val Glu Asp 210 Ser Lys Leu Cys Lys Ile Cys Tyr Val Glu Glu Cys Ile Val Cys Phe 235 Val Pro Cys Gly His Val Val Ala Cys Ala Lys Cys Ala Leu Ser Val 250 Asp Lys Cys Pro Met Cys Arg Lys Ile Val Thr Ser Val Leu Lys Val 270 260 Tyr Phe Ser

(2) INFORMATION FOR SEQ ID NO:13:

275

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 498 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met Thr Glu Leu Gly Met Glu Leu Glu Ser Val Arg Leu Ala Thr Phe 10 10 15

Gly Glu Trp Pro Leu Asn Ala Pro Val Ser Ala Glu Asp Leu Val Ala 20 25 30

Asn Gly Phe Phe Ala Thr Gly Lys Trp Leu Glu Ala Glu Cys His Phe 35 40 45

Cys His Val Arg Ile Asp Arg Trp Glu Tyr Gly Asp Gln Val Ala Glu 50 60

Arg His Arg Arg Ser Ser Pro Ile Cys Ser Met Val Leu Ala Pro Asn 70 75 80

His Cys Gly Asn Val Pro Arg Ser Gln Glu Ser Asp Asn Glu Gly Asn 85 90 95

Ser Val Val Asp Ser Pro Glu Ser Cys Ser Cys Pro Asp Leu Leu Leu 100 Glu Ala Asn Arg Leu Val Thr Phe Lys Asp Trp Pro Asn Pro Asn Ile 120 Thr Pro Gln Ala Leu Ala Lys Ala Gly Phe Tyr Tyr Leu Asn Arg Leu Asp His Val Lys Cys Val Trp Cys Asn Gly Val Ile Ala Lys Trp Glu 145 Lys Asn Asp Asn Ala Phe Glu Glu His Lys Arg Phe Phe Pro Gln Cys Pro Arg Val Gln Met Gly Pro Leu Ile Glu Phe Ala Thr Gly Lys Asn 185 Leu Asp Glu Leu Gly Ile Gln Pro Thr Thr Leu Pro Leu Arg Pro Lys 200 Tyr Ala Cys Val Asp Ala Arg Leu Arg Thr Phe Thr Asp Trp Pro Ile Ser Asn Ile Gln Pro Ala Ser Ala Leu Ala Gln Ala Gly Leu Tyr Tyr 230 235 Gln Lys Ile Gly Asp Gln Val Arg Cys Phe His Cys Asn Ile Gly Leu Arg Ser Trp Gln Lys Glu Asp Glu Pro Trp Phe Glu His Ala Lys Trp Ser Pro Lys Cys Gln Phe Val Leu Leu Ala Lys Gly Pro Ala Tyr Val 280 Ser Glu Val Leu Ala Thr Thr Ala Ala Asn Ala Ser Ser Gln Pro Ala 295 290 Thr Ala Pro Ala Pro Thr Leu Gln Ala Asp Val Leu Met Asp Glu Ala 310 315 Pro Ala Lys Glu Ala Leu Thr Leu Gly Ile Asp Gly Gly Val Val Arg Asn Ala Ile Gln Arg Lys Leu Leu Ser Ser Gly Cys Ala Phe Ser Thr Leu Asp Glu Leu Leu His Asp Ile Phe Asp Asp Ala Gly Ala Gly Ala 360 Ala Leu Glu Val Arg Glu Pro Pro Glu Pro Ser Ala Pro Phe Ile Glu 375 Pro Cys Gln Ala Thr Thr Ser Lys Ala Ala Ser Val Pro Ile Pro Val 390 385 Ala Asp Ser Ile Pro Ala Lys Pro Gln Ala Ala Glu Ala Val Ser Asn 410 Ile Ser Lys Ile Thr Asp Glu Ile Gln Lys Met Ser Val Ser Thr Pro 420 425

Asn Gly Asn Leu Ser Leu Glu Glu Glu Asn Arg Gln Leu Lys Asp Ala 435 440 445

Arg Leu Cys Lys Val Cys Leu Asp Glu Glu Val Gly Val Val Phe Leu 450 460

Pro Cys Gly His Leu Ala Thr Cys Asn Gln Cys Ala Pro Ser Val Ala 465 470 475 480

Asn Cys Pro Met Cys Arg Ala Asp Ile Lys Gly Phe Val Arg Thr Phe 485 490 495

Leu Ser

- (2) INFORMATION FOR SEQ ID NO:14:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 67 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Glu Glu Val Arg Leu Asn Thr Phe Glu Lys Trp Pro Val Ser Phe Leu 1 5 10 15

Ser Pro Glu Thr Met Ala Lys Asn Gly Phe Tyr Tyr Leu Gly Arg Ser 20 25 30

Asp Glu Val Arg Cys Ala Phe Cys Lys Val Glu Ile Met Arg Trp Lys 35 40 45

Glu Gly Glu Asp Pro Ala Ala Asp His Lys Lys Trp Ala Pro Gln Cys 50 55 60

Pro Phe Val

- (2) INFORMATION FOR SEQ ID NO:15:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 67 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:

Glu Ala Asn Arg Leu Val Thr Phe Lys Asp Trp Pro Asn Pro Asn Ile 1 5 10 15 Thr Pro Gln Ala Leu Ala Lys Ala Gly Phe Tyr Tyr Leu Asn Arg Leu 20 25 30

Asp His Val Lys Cys Val Trp Cys Asn Gly Val Ile Ala Lys Trp Glu 35 40 45

Lys Asn Asp Asn Ala Phe Glu Glu His Lys Arg Phe Phe Pro Gln Cys 50 55 60

Pro Arg Val 65

- (2) INFORMATION FOR SEQ ID NO:16:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:
  - Glu Phe Asn Arg Leu Lys Thr Phe Ala Asn Phe Pro Ser Ser Pro 10 15
  - Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu 20 25 30
  - Gly Asp Thr Val Gln Cys Phe Ser Cys His Ala Ala Ile Asp Arg Trp 35 40 45
  - Gln Tyr Gly Asp Ser Ala Val Gly Arg His Arg Arg Ile Ser Pro Asn 50 55 60

Cys Arg Phe Ile

- (2) INFORMATION FOR SEQ ID NO:17:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:17:
  - Glu Phe Asn Arg Leu Lys Thr Phe Ala Asn Phe Pro Ser Gly Ser Pro 1 5 10 15
  - Val Ser Ala Ser Thr Leu Ala Arg Ala Gly Phe Leu Tyr Thr Gly Glu 20 25 30

Gly Asp Thr Val Arg Cys Phe Ser Cys His Ala Ala Val Asp Arg Trp
35 40 45

Gln Tyr Gly Asp Ser Ala Val Gly Arg His Arg Lys Val Ser Pro Asn 50 55 60

Cys Arg Phe Ile

- (2) INFORMATION FOR SEQ ID NO:18:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:18:

Glu Leu Tyr Arg Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro
1 5 10 15

Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val 20 25 30

Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp 35 40 45

Lys Arg Gly Asp Ser Pro Thr Glu Lys His Lys Leu Tyr Pro Ser 50 60

Cys Arg Phe Val 65

- (2) INFORMATION FOR SEQ ID NO:19:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

Glu Leu Tyr Arg Met Ser Thr Tyr Ser Thr Phe Pro Ala Gly Val Pro 1 5 10 15

Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val 20 25 30

Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp 35 40 45

Lys Leu Gly Asp Ser Pro Ile Gln Lys His Lys Gln Leu Tyr Pro Ser 50 60

Cys Ser Phe Ile

- (2) INFORMATION FOR SEQ ID NO:20:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr Ala His 1 5 10 15

Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr Gly Ala 20 25 30

Asp Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp 35 40 45

Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn 50 55 60

Cys Phe Phe Val

- (2) INFORMATION FOR SEQ ID NO:21:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Glu Glu Ala Arg Leu Lys Ser Phe Gln Asn Trp Pro Asp Tyr Ala His
1 10 15

Leu Thr Pro Arg Glu Leu Ala Ser Ala Gly Leu Tyr Tyr Thr Gly Ile 20 25 30

Gly Asp Gln Val Gln Cys Phe Cys Cys Gly Gly Lys Leu Lys Asn Trp 35 40 45

Glu Pro Cys Asp Arg Ala Trp Ser Glu His Arg Arg His Phe Pro Asn 50 55 60

Cys Phe Phe Val 65

- (2) INFORMATION FOR SEQ ID NO:22:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 67 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:22:

Glu Asn Ala Arg Leu Leu Thr Phe Gln Thr Trp Pro Leu Thr Phe Leu 1 5 10 15

Ser Pro Thr Asp Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly 20 25 30

Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu 35 40 45

Pro Lys Asp Asn Ala Met Ser Glu His Leu Arg His Phe Pro Lys Cys 50 60

Pro Phe Ile 65

- (2) INFORMATION FOR SEQ ID NO:23:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 67 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

Glu Glu Ala Arg Phe Leu Thr Tyr His Met Trp Pro Leu Thr Phe Leu
1 5 10 15

Ser Pro Ser Glu Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly 20 25 30

Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu

Pro Lys Asp Asp Ala Met Ser Glu His Arg Arg His Phe Pro Asn Cys 50 55 60

Pro Phe Leu 65

#### (2) INFORMATION FOR SEQ ID NO:24:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 66 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

Tyr Glu Ala Arg Ile Val Thr Phe Gly Thr Trp Ile Tyr Ser Val Asn 1 5 10 15

Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp 20 25 30

Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro 35 40 45

Ser Glu Asp Pro Trp Asp Gln His Ala Lys Cys Tyr Pro Gly Cys Lys 50 60

Tyr Leu 65

- (2) INFORMATION FOR SEQ ID NO:25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 66 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:25:

Tyr Glu Ala Arg Ile Phe Thr Phe Gly Thr Trp Ile Tyr Ser Val Asn 1 5 10 15

Lys Glu Gln Leu Ala Arg Ala Gly Phe Tyr Ala Leu Gly Glu Gly Asp 20 25 30

Lys Val Lys Cys Phe His Cys Gly Gly Gly Leu Thr Asp Trp Lys Pro 35 40 45

Ser Glu Asp Pro Trp Glu Gln His Ala Lys Trp Tyr Pro Gly Cys Lys 50 55 60

Tyr Leu 65

(2) INFORMATION FOR SEQ ID NO:26:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 68 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:26:

His Ala Ala Arg Phe Lys Thr Phe Phe Asn Trp Pro Ser Ser Val Leu 1 10 15

Val Asn Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Asn 20 25 30

Ser Asp Asp Val Lys Cys Phe Cys Cys Asp Gly Gly Leu Arg Cys Trp
35 40 45

Glu Ser Gly Asp Asp Pro Trp Val Gln His Ala Lys Trp Phe Pro Arg 50 55 60

Cys Glu Tyr Leu 65

- (2) INFORMATION FOR SEQ ID NO:27:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

His Ala Ala Arg Met Arg Thr Phe Met Tyr Trp Pro Ser Ser Val Pro 1 5 10 15

Val Gln Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Gly Arg 20 25 30

Asn Asp Asp Val Lys Cys Phe Gly Cys Asp Gly Gly Leu Arg Cys Trp 35 40 45

Glu Ser Gly Asp Asp Pro Trp Val Glu His Ala Lys Trp Phe Pro Arg
50 55 60

Cys Glu Phe Leu 65

- (2) INFORMATION FOR SEQ ID NO:28:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids

- (B) TYPE: amino acid
- (C) STRANDEDNESS: not relevant
- (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:
- Glu Ala Ala Arg Leu Arg Thr Phe Ala Glu Trp Pro Arg Gly Leu Lys
  1 10 15
- Gln Arg Pro Glu Glu Leu Ala Glu Ala Gly Phe Phe Tyr Thr Gly Gln 20 25 30
- Gly Asp Lys Thr Arg Cys Phe Cys Cys Asp Gly Gly Leu Lys Asp Trp 35 40 45
- Glu Pro Asp Asp Ala Pro Trp Gln Gln His Ala Arg Trp Tyr Asp Arg 50 55 60

Cys Glu Tyr Val 65

- (2) INFORMATION FOR SEQ ID NO:29:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:29:
  - Glu Ala Ala Arg Val Lys Ser Phe His Asn Trp Pro Arg Cys Met Lys 1 5 10 15
  - Gln Arg Pro Glu Gln Met Ala Asp Ala Gly Phe Phe Tyr Thr Gly Tyr 20 25 30
  - Gly Asp Asn Thr Lys Cys Phe Tyr Cys Asp Gly Gly Leu Lys Asp Trp 35 40 45
  - Glu Pro Glu Asp Val Pro Trp Glu Gln His Val Arg Trp Phe Asp Arg 50 55 60

Cys Ala Tyr Val

- (2) INFORMATION FOR SEQ ID NO:30:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 68 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant

#### (D) TOPOLOGY: both

#### (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

Val Asp Ala Arg Leu Arg Thr Phe Thr Asp Trp Pro Ile Ser Asn Ile 1 5 10 15

Gln Pro Ala Ser Ala Leu Ala Gln Ala Gly Leu Tyr Tyr Gln Lys Ile 20 25 30

Gly Asp Gln Val Arg Cys Phe His Cys Asn Ile Gly Leu Arg Ser Trp 35 40 45

Gln Lys Glu Asp Glu Pro Trp Phe Glu His Ala Lys Trp Ser Pro Lys 50 55 60

Cys Gln Phe Val

#### (2) INFORMATION FOR SEQ ID NO:31:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 66 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

Glu Ser Val Arg Leu Ala Thr Phe Gly Glu Trp Pro Leu Asn Ala Pro 1 5 10 15

Val Ser Ala Glu Asp Leu Val Ala Asn Gly Phe Phe Gly Thr Trp Met 20 25 30

Glu Ala Glu Cys Asp Phe Cys His Val Arg Ile Asp Arg Trp Glu Tyr 35 40 45

Gly Asp Leu Val Ala Glu Arg His Arg Arg Ser Ser Pro Ile Cys Ser 50 60

Met Val

#### (2) INFORMATION FOR SEQ ID NO:32:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 46 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both

## (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:

Glu Gln Leu Arg Arg Leu Gln Glu Glu Arg Thr Cys Lys Val Cys Met
1 5 10 15

Asp Lys Glu Val Ser Val Val Phe Ile Pro Cys Gly His Leu Val Val 20 25 30

Cys Gln Glu Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys 35 40 45

- (2) INFORMATION FOR SEQ ID NO:33:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 46 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:

Glu Gln Leu Arg Arg Leu Pro Glu Glu Arg Thr Cys Lys Val Cys Met
1 10 15

Asp Lys Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val 20 25 30

Cys Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys 35 40 45

- (2) INFORMATION FOR SEQ ID NO:34:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 46 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:

Glu Gln Leu Arg Arg Leu Gln Glu Glu Lys Leu Ser Lys Ile Cys Met
1 5 10 15

Asp Arg Asn Ile Ala Ile Val Phe Phe Pro Cys Gly His Leu Ala Thr

Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys Pro Met Cys

#### (2) INFORMATION FOR SEQ ID NO:35:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 46 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

Glu Gln Leu Arg Arg Leu Gln Glu Glu Lys Leu Cys Lys Ile Cys Met
1 5 10 15

Asp Arg Asn Ile Ala Ile Val Phe Val Pro Cys Gly His Leu Val Thr 20 25 30

Cys Lys Gln Cys Ala Glu Ala Val Asp Lys Cys Pro Met Cys 35 40 45

- (2) INFORMATION FOR SEQ ID NO:36:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 46 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

Glu Glu Asn Arg Gln Leu Lys Asp Ala Arg Leu Cys Lys Val Cys Leu 1 5 10 15

Asp Glu Glu Val Gly Val Val Phe Leu Pro Cys Gly His Leu Ala Thr 20 25 30

Cys Asn Gln Cys Ala Pro Ser Val Ala Asn Cys Pro Met Cys 35 40 45

- (2) INFORMATION FOR SEQ ID NO:37:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 46 amino acids
    - (B) TYPE: amino acid
    - (C) STRANDEDNESS: not relevant
    - (D) TOPOLOGY: both
  - (ii) MOLECULE TYPE: protein

(xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:37:
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Glu Lys Glu Pro Gln Val Glu Asp Ser Lys Leu Cys Lys Ile Cys Tyr
1 5 10 15

Val Glu Glu Cys Ile Val Cys Phe Val Pro Cys Gly His Val Val Ala
20 25 30

Cys Ala Lys Cys Ala Leu Ser Val Asp Lys Cys Pro Met Cys 35 40 45

#### (2) INFORMATION FOR SEQ ID NO:38:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 46 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: both
- (ii) MOLECULE TYPE: protein

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:38:

Ala Val Glu Ala Glu Val Ala Asp Asp Arg Leu Cys Lys Ile Cys Leu
1 5 10 15

Gly Ala Glu Lys Thr Val Cys Phe Val Pro Cys Gly His Val Val Ala 20 25 30

Cys Gly Lys Cys Ala Ala Gly Val Thr Thr Cys Pro Val Cys 35 40 45

#### (2) INFORMATION FOR SEQ ID NO:39:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2474 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:39:

60	CAGAGCCTAG	TGCTGGCGTT	CAGAGGTCAT	CCCCGGAGAT	AGACCTACAC	GAATTCCGGG
120	ACAAAACTAC	GAAGCCATGC	AAACCGACCA	CCTAGCAGTA	GCGGTATCAG	GAAGTGGGCT
180	CATGGTTCAA	TCACCATGAA	CCCTGTCATC	CCCTTCCCCT	AAAGACTTGT	ATCCCCAGAG
240	GAAGTATGAC	CCTTTGAGTT	AGTGCTGACA	GCTGATGAAG	TTCTAGCCAA	GACAGCGCCT
300	AGTTCCTGTG	TTCCCAGGGG	TATTCAGCTT	ATTGTCCACG	AGCTGTACCG	TTTTCCTGTG
360	CAAGGTCAAG	GTGCCAATGA	TACTACACTG	TGCTGGCTTT	GTCTGGCTCG	TCAGAAAGGA

TGCTTCTGCT	GTGGCCTGAT	GCTAGACAAC	TGGAAACAAG	GGGACAGTCC	CATGGAGAAG	420
CACAGAAAGT	TGTACCCCAG	CTGCAACTTT	GTACAGACTT	TGAATCCAGC	CAACAGTCTG	480
GAAGCTAGTC	CTCGGCCTTC	TCTTCCTTCC	ACGGCGATGA	GCACCATGCC	TTTGAGCTTT	540
GCAAGTTCTG	AGAATACTGG	CTATTTCAGT	GGCTCTTACT	CGAGCTTTCC	CTCAGACCCT	600
GTGAACTTCC	GAGCAAATCA	AGATTGTCCT	GCTTTGAGCA	CAAGTCCCTA	CCACTTTGCA	660
ATGAACACAG	AGAAGGCCAG	ATTACTCACC	TATGAAACAT	GGCCATTGTC	TTTTCTGTCA	720
CCAGCAAAGC	TGGCCAAAGC	AGGCTTCTAC	TACATAGGAC	CTGGAGATAG	AGTGGCCTGC	780
TTTGCGTGCG	ATGGGAAACT	GAGCAACTGG	GAACGTAAGG	ATGATGCTAT	GTCAGAGCAC	840
CAGAGGCATT	TCCCCAGCTG	TCCGTTCTTA	AAAGACTTGG	GTCAGTCTGC	TTCGAGATAC	900
ACTGTCTCTA	ACCTGAGCAT	GCAGACACAC	GCAGCCCGTA	TTAGAACATT	CTCTAACTGG	960
CCTTCTAGTG	CACTAGTTCA	TTCCCAGGAA	CTTGCAAGTG	CGGGCTTTTA	TTATACAGGA	1020
CACAGTGATG	ATGTCAAGTG	TTTATGCTGT	GATGGTGGGC	TGAGGTGCTG	GGAATCTGGA	1080
GATGACCCCT	GGGTGGAACA	TGCCAAGTGG	TTTCCAAGGT	GTGAGTACTT	GCTCAGAATC	1140
AAAGGCCAAG	AATTTGTCAG	CCAAGTTCAA	GCTGGCTATC	CTCATCTACT	TGAGCAGCTA	1200
TTATCTACGT	CAGACTCCCC	AGAAGATGAG	AATGCAGACG	CAGCAATCGT	GCATTTTGGC	1260
CCTGGAGAAA	GTTCGGAAGA	TGTCGTCATG	ATGAGCACGC	CTGTGGTTAA	AGCAGCCTTG	1320
GAAATGGGCT	TCAGTAGGAG	CCTGGTGAGA	CAGACGGTTC	AGTGGCAGAT	CCTGGCCACT	1380
GGTGAGAACT	ACAGGACCGT	CAGTGACCTC	GTTATAGGCT	TACTCGATGC	AGAAGACGAG	1440
ATGAGAGAGG	AGCAGATGGA	GCAGGCGGCC	GAGGAGGAGG	AGTCAGATGA	TCTAGCACTA	1500
ATCCGGAAGA	ACAAAATGGT	GCTTTTCCAA	CATTTGACGT	GTGTGACACC	AATGCTGTAT	1560
TGCCTCCTAA	GTGCAAGGGC	CATCACTGAA	CAGGAGTGCA	ATGCTGTGAA	ACAGAAACCA	1620
CACACCTTAC	AAGCAAGCAC	ACTGATTGAT	ACTGTGTTAG	CAAAAGGAAA	CACTGCAGCA	1680
ACCTCATTCA	GAAACTCCCT	TCGGGAAATT	GACCCTGCGT	TATACAGAGA	TATATTTGTG	1740
CAACAGGACA	TTAGGAGTCT	TCCCACAGAT	GACATTGCAG	CTCTACCAAT	GGAAGAACAG	1800
TTGCGGCCCC	TCCCGGAGGA	CAGAATGTGT	AAAGTGTGTA	TGGACCGAGA	GGTATCCATC	1860
GTGTTCATTC	CCTGTGGCCA	TCTGGTCGTG	TGCAAAGACT	GCGCTCCCTC	TCTGAGGAAG	1920
TGTCCCATCT	GTAGAGGGAC	CATCAAGGGC	ACAGTGCGCA	CATTTCTCTC	CTGAACAAGA	1980
CTAATGGTCC	ATGGCTGCAA	CTTCAGCCAG	GAGGAAGTTC	ACTGTCACTC	CCAGTTCCAT	2040
TCGGAACTTG	AGGCCAGCCT	GGATAGCACG	AGACACCGCC	AAACACACAA	ATATAAACAT	2100
GAAAAACTTT	TGTCTGAAGT	CAAGAATGAA	TGAATTACTT	ATATAATAAT	TTTAATTGGT	2160
TTCCTTAAAA	GTGCTATTTG	TTCCCAACTC	AGAAAATTGT	TTTCTGTAAA	CATATTTACA	2220

TACTACCTGC	ATCTAAAGTA	TTCATATATT	CATATATTCA	GATGTCATGA	GAGAGGGTTT	2280
TGTTCTTGTT	CCTGAAAAGC	TGGTTTATCA	TCTGATCAGC	ATATACTGCG	CAACGGGCAG	2340
GGCTAGAATC	CATGAACCAA	GCTGCAAAGA	TCTCACGCTA	AATAAGGCGG	AAAGATTTGG	2400
AGAAACGAAA	GGAAATTCTT	TCCTGTCCAA	TGTATACTCT	TCAGACTAAT	GACCTCTTCC	2460
TATCAAGCCT	TCTA					2474

# (2) INFORMATION FOR SEQ ID NO:40:

- (i) SEQUENCE CHARACTERISTICS:

  - (A) LENGTH: 602 amino acids
    (B) TYPE: amino acid
    (C) STRANDEDNESS: not relevant
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:40:
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(XI)	SEQU	ENCE	, DES	CKI	TION	. JL	, Q . L	, 110.	. 40.						
Met 1	Asn	Met	Val	Gln 5	Asp	Ser	Ala	Phe	Leu 10	Ala	Lys	Leu	Met	Lys 15	Ser
Ala	Asp	Thr	Phe 20	Glu	Leu	Lys	Tyr	Asp 25	Phe	Ser	Cys	Glu	Leu 30	Tyr	Arg
Leu	Ser	Thr 35	Tyr	Ser	Ala	Phe	Pro 40	Arg	Gly	Val	Pro	Val 45	Ser	Glu	Arg
Ser	Leu 50	Ala	Arg	Ala	Gly	Phe 55	Tyr	Tyr	Thr	Gly	Ala 60	Asn	Asp	Lys	Val
Lys 65	Cys	Phe	Cys	Cys	Gly 70	Leu	Met	Leu	Asp	Asn 75	Trp	Lys	Gln	Gly	Asp 80
Ser	Pro	Met	Glu	Lys 85	His	Arg	Lys	Leu	Tyr 90	Pro	Ser	Cys	Asn	Phe 95	Val
Gln	Thr	Leu	Asn 100	Pro	Ala	Asn	Ser	Leu 105	Glu	Ala	Ser	Pro	Arg 110	Pro	Ser
Leu	Pro	Ser 115	Thr	Ala	Met	Ser	Thr 120	Met	Pro	Leu	Ser	Phe 125	Ala	Ser	Ser
Glu	Asn 130	Thr	Gly	Tyr	Phe	Ser 135	Gly	Ser	Tyr	Ser	Ser 140	Phe	Pro	Ser	Asp
Pro 145	Val	Asn	Phe	Arg	Ala 150	Asn	Gln	Asp	Cys	Pro 155	Ala	Leu	Ser	Thr	Ser 160
Pro	Tyr	His	Phe	Ala 165	Met	Asn	Thr	Glu	Lys 170	Ala	Arg	Leu	Leu	Thr 175	Tyr
Glu	Thr	Trp	Pro 180	Leu	Ser	Phe	Leu	Ser 185	Pro	Ala	Lys	Leu	Ala 190	Lys	Ala
Gly	Phe	Tyr	Tyr	Ile	Gly	Pro	Gly	Asp	Arg	Val	Ala	Cys	Phe	Ala	Cys

		195					200					205			
Asp	Gly 210	Lys	Leu	Ser	Asn	Trp 215	Glu	Arg	Lys	Asp	Asp 220	Ala	Met	Ser	Glu
His 225	Gln	Arg	His	Phe	Pro 230	Ser	Cys	Pro	Phe	Leu 235	Lys	Asp	Leu	Gly	Gln 240
Ser	Ala	Ser	Arg	Tyr 245	Thr	Val	Ser	Asn	Leu 250	Ser	Met	Gln	Thr	His 255	Ala
Ala	Arg	Ile	Arg 260	Thr	Phe	Ser	Asn	Trp 265	Pro	Ser	Ser	Ala	Leu 270	Val	His
Ser	Gln	Glu 275	Leu	Ala	Ser	Ala	Gly 280	Phe	Tyr	Tyr	Thr	Gly 285	His	Ser	Asp
Asp	Val 290	Lys	Cys	Leu	Cys	Cys 295	Asp	Gly	Gly	Leu	Arg 300	Cys	Trp	Glu	Ser
Gly 305	Asp	Asp	Pro	Trp	Val 310	Glu	His	Ala	Lys	Trp 315	Phe	Pro	Arg	Cys	Glu 320
Tyr	Leu	Leu	Arg	Ile 325	Lys	Gly	Gln	Glu	Phe 330	Val	Ser	Gln	Val	Gln 335	Ala
Gly	Tyr	Pro	His 340	Leu	Leu	Glu	Gln	Leu 345	Leu	Ser	Thr	Ser	Asp 350	Ser	Pro
Glu	Asp	Glu 355	Asn	Ala	Asp	Ala	Ala 360	Ile	Val	His	Phe	Gly 365	Pro	Gly	Glu
Ser	Ser 370	Glu	Asp	Val	Val	Met 375	Met	Ser	Thr	Pro	Val 380	Val	Lys	Ala	Ala
Leu 385	Glu	Met	Gly	Phe	Ser 390	Arg	Ser	Leu	Val	Arg 395	Gln	Thr	Val	Gln	Trp 400
Gln	Ile	Leu	Ala	Thr 405	Gly	Glu	Asn	Tyr	Arg 410	Thr	Val	Ser	Asp	Leu 415	Val
Ile	Gly	Leu	Leu 420	Asp	Ala	Glu	Asp	Glu 425	Met	Arg	Glu	Glu	Gln 430	Met	Glu
Gln	Ala	Ala 435	Glu	Glu	Glu	Glu	Ser 440	Asp	Asp	Leu	Ala	Leu 445	Ile	Arg	Lys
Asn	Lys 450	Met	Val	Leu	Phe	Gln 455	His	Leu	Thr	Cys	Val 460	Thr	Pro	Met	Leu
Tyr 465	Cys	Leu	Leu	Ser	Ala 470	Arg	Ala	Ile	Thr	Glu 475	Gln	Glu	Cys	Asn	Ala 480
Val	Lys	Gln	Lys	Pro 485	His	Thr	Leu	Gln	Ala 490	Ser	Thr	Leu	Ile	Asp 495	Thr
Val	Leu	Ala	Lys 500	Gly	Asn	Thr	Ala	Ala 505	Thr	Ser	Phe	Arg	Asn 510	Ser	Leu
Arg	Glu	Ile 515	Asp	Pro	Ala	Leu	Tyr 520	Arg	Asp	Ile	Phe	Val 525	Gln	Gln	Asp

Ile Arg Ser Leu Pro Thr Asp Asp Ile Ala Ala Leu Pro Met Glu Glu 530 540

Gln Leu Arg Pro Leu Pro Glu Asp Arg Met Cys Lys Val Cys Met Asp 545 550 555 560

Arg Glu Val Ser Ile Val Phe Ile Pro Cys Gly His Leu Val Val Cys 565 570 575

Lys Asp Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys Arg Gly Thr 580 585 590

Ile Lys Gly Thr Val Arg Thr Phe Leu Ser 595 600

#### (2) INFORMATION FOR SEQ ID NO:41:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 2416 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

60	TAAGCGGTCA	TCTGGAAGTT	AGAAACTTCA	CCAAGTGGTG	GATCTATTGT	CTGTGGTGGA
120	AGGTACCTTA	GACTCGCCCA	GTCTCCCAGA	GGACAAAACT	TACTACTCAT	GAAATACTAT
180	GGACAAAGGA	TTGTCAAATT	GAGCACAATC	TAATGGAGAA	CTTAAACGTA	CACCCAAAAA
240	CTACATATTC	TACCGAATGT	GTGTGAACTC	TTGACTTTTC	AAAATGAAGT	GAGCGAAGAA
300	GCTTTTATTA	GCTCGTGCTG	GAGGAGTCTG	CTGTCTCAGA	AGGGGAGTTC	AGCTTTTCCC
360	ATAACTGGAA	CTGATGTTGG	CTGCTGTGGC	TCAAGTGCTT	AATGACAAAG	TACAGGTGTG
420	GCTTTGTACA	CCCAGCTGCA	ACAGTTCTAT	AAAAGCACAG	AGTCCTGTTG	ACAAGGGGAC
480	TGAAAAGTAG	ATGTCTCCTG	ATCTAAGAAT	TGCAGTCTCC	TCAGCCAGTC	GACTCTGCTT
540	GCTCTAGCCC	TCCAACCTGT	TGGCATTCAC	TGGAACGAGG	TCGTCACCTC	ATTTGCACAT
600	GCTATGCCAT	GATCCCTGCA	ATCAAGGATG	AAGACTTCTC	AGAGCAGTGG	TCTTAATTCT
660	TTCTGTCACC	CCTTTAAGTT	CAGTATGTGG	TTCTTACTTA	GAGGCCAGAT	GAGTACAGAA
720	TGGCCTGTTT	GGAGACAGGG	CATAGGGCCT	GCTTCTATTA	GCCAGAGCTG	AGCAGAGCTG
780	CAGAGCACCG	TATGCTATGT	ACCAAAGGAT	GCAACTGGGA	GGGAAACTGA	TGCCTGTGGT
840	GGTTTAGTAT	GAAACACAGA	AAATACTTCA	CATTTCTGGA	CCCCACTGTC	CAGACATTTT
900	ACTGGCCACC	ACATTTCTGT	TCGATTGAGG	CACACTCTGC	AGTATGCAGA	ATCAAATCTA
960	TGGATCGCAA	TTCTATTACG	AAGTGCTGGA	AGCAGCTTGC	GTTCAGCCCG	TAGTGTTCCT



#### (2) INFORMATION FOR SEQ ID NO:42:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 591 amino acids
  - (B) TYPE: amino acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

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## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:

Met Glu Lys Ser Thr Ile Leu Ser Asn Trp Thr Lys Glu Ser Glu Glu Lys Met Lys Phe Asp Phe Ser Cys Glu Leu Tyr Arg Met Ser Thr Tyr Ser Ala Phe Pro Arg Gly Val Pro Val Ser Glu Arg Ser Leu Ala Arg Ala Gly Phe Tyr Tyr Thr Gly Val Asn Asp Lys Val Lys Cys Phe Cys Cys Gly Leu Met Leu Asp Asn Trp Lys Gln Gly Asp Ser Pro Val Glu Lys His Arg Gln Phe Tyr Pro Ser Cys Ser Phe Val Gln Thr Leu Leu Ser Ala Ser Leu Gln Ser Pro Ser Lys Asn Met Ser Pro Val Lys Ser Arg Phe Ala His Ser Ser Pro Leu Glu Arg Gly Gly Ile His Ser Asn Leu Cys Ser Ser Pro Leu Asn Ser Arg Ala Val Glu Asp Phe Ser Ser Arg Met Asp Pro Cys Ser Tyr Ala Met Ser Thr Glu Glu Ala Arg Phe 145 150 155 Leu Thr Tyr Ser Met Trp Pro Leu Ser Phe Leu Ser Pro Ala Glu Leu Ala Arg Ala Gly Phe Tyr Tyr Ile Gly Pro Gly Asp Arg Val Ala Cys Phe Ala Cys Gly Gly Lys Leu Ser Asn Trp Glu Pro Lys Asp Tyr Ala Met Ser Glu His Arg Arg His Phe Pro His Cys Pro Phe Leu Glu Asn 215 Thr Ser Glu Thr Gln Arg Phe Ser Ile Ser Asn Leu Ser Met Gln Thr 230 His Ser Ala Arg Leu Arg Thr Phe Leu Tyr Trp Pro Pro Ser Val Pro 245 Val Gln Pro Glu Gln Leu Ala Ser Ala Gly Phe Tyr Tyr Val Asp Arg Asn Asp Asp Val Lys Cys Leu Cys Cys Asp Gly Gly Leu Arg Cys Trp Glu Pro Gly Asp Asp Pro Trp Ile Glu His Ala Lys Trp Phe Pro Arg 290 295 Cys Glu Phe Leu Ile Arg Met Lys Gly Gln Glu Phe Val Asp Glu Ile

315

310

Gln Ala Arg Tyr Pro His Leu Leu Glu Gln Leu Leu Ser Thr Ser Asp 330 Thr Pro Gly Glu Glu Asn Ala Asp Pro Thr Glu Thr Val Val His Phe Gly Pro Gly Glu Ser Ser Lys Asp Val Val Met Met Ser Thr Pro Val 360 Val Lys Ala Ala Leu Glu Met Gly Phe Ser Arg Ser Leu Val Arg Gln Thr Val Gln Arg Gln Ile Leu Ala Thr Gly Glu Asn Tyr Arg Thr Val 385 390 395 Asn Asp Ile Val Ser Val Leu Leu Asn Ala Glu Asp Glu Arg Arg Glu Glu Glu Lys Glu Arg Gln Thr Glu Glu Met Ala Ser Gly Asp Leu Ser Leu Ile Arg Lys Asn Arg Met Ala Leu Phe Gln Gln Leu Thr His Val Leu Pro Ile Leu Asp Asn Leu Leu Glu Ala Ser Val Ile Thr Lys Gln Glu His Asp Ile Ile Arg Gln Lys Thr Gln Ile Pro Leu Gln Ala Arg Glu Leu Ile Asp Thr Val Leu Val Lys Gly Asn Ala Ala Asn Ile Phe Lys Asn Ser Leu Lys Gly Ile Asp Ser Thr Leu Tyr Glu Asn Leu Phe Val Glu Lys Asn Met Lys Tyr Ile Pro Thr Glu Asp Val Ser Gly 515 520 525 Leu Ser Leu Glu Glu Gln Leu Arg Arg Leu Gln Glu Glu Arg Thr Cys Lys Val Cys Met Asp Arg Glu Val Ser Ile Val Phe Ile Pro Cys Gly 550 555 His Leu Val Val Cys Gln Glu Cys Ala Pro Ser Leu Arg Lys Cys Pro Ile Cys Arg Gly Thr Ile Lys Gly Thr Val Arg Thr Phe Leu Ser 580 585